substituted cyclic hydrocarbon group; R1 is hydrogen, optionally substituted hydrocarbyl, an optionally substituted heterocyclic group, or acyl; R2 is optionally substituted amino; D is a free valency or a divalent group; E is CO, CON(Ra), COO, N(Ra)CON(Rb), N(Ra)SO2, N(Ra), O, S, SO, SO2; G is a free valency or a divalent group; L is a free valency, an optionally substituted divalent hydrocarbon group which may be interrupted by O or S, or the like; X is oxygen, optionally oxidized sulfur, optionally substituted nitrogen, or an optionally substituted divalent hydrocarbon group; Y is two hydrogen atoms, oxygen, or sulfur; and the dotted line indicates that R2 and an atom on ring B may together form a ring] and salts are prepd. and tested as somatostatin receptor regulators. Thus, the title compd. II was prepd. in treatment or prevention of diabetes and obesity.

IT 32897-26-0

RL: RCT (Reactant)

(prepn. of arom. amine derivs. and agents contg. the same as somatostatin receptor regulators)

RN 32897-26-0 CAPLUS

Urea, N-[3-(dimethylamino)propyl]-N'-ethyl- (9CI) (CA INDEX NAME) CN

0 EtNH-C-NH-(CH₂)₃-NMe₂

REFERENCE COUNT:

REFERENCE(S):

- (1) Eastman Kodak Company; DE 2855697 Al CAPLUS
- (2) Eastman Kodak Company; JP 54145135 A CAPLUS
- (3) Eastman Kodak Company; JP 54145135 A CAPLUS
- (4) Eastman Kodak Company; GB 2010818 A 1979 CAPLUS
- (5) Fuji Photo Film Co Ltd; JP 61233741 A 1986 CAPLUS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L12 ANSWER 6 OF 15 CAPLUS COPYRIGHT 2001 ACS ACCESSION NUMBER:

DOCUMENT NUMBER:

1997:626481 CAPLUS

TITLE:

127:262918

Synthesis of carbohydrate-containing dendrimers. 5.

Preparation of dendrimers using unprotected

carbohydrates

AUTHOR(S):

Jayaraman, Narayanaswamy; Stoddart, J. Fraser

CORPORATE SOURCE:

Sch. Chem., Univ. Birmingham, Birmingham, B15 2TT, UK

SOURCE:

Tetrahedron Lett. (1997), 38(38), 6767-6770

CODEN: TELEAY; ISSN: 0040-4039

PUBLISHER:

Elsevier Journal

DOCUMENT TYPE:

English LANGUAGE:

Carbohydrate-contg. dendrimers have been prepd. using completely unprotected carbohydrates employing a convergent growth approach. facile syntheses of lower generation dendrimers, using the amide bond forming methodol., opens up the possibility of obtaining densely-packed glycodendrimers without the need to resort to protecting group manipulations on the saccharide residues.

32897-26-0 ΙT

RL: RCT (Reactant)

(prepn. of dendrimers using unprotected carbohydrates)

RN 32897-26-0 CAPLUS CN Urea, N-[3-(dimethylamino)propyl]-N'-ethyl- (9CI) (CA INDEX NAME)

O || EtNH-C-NH-(CH₂)3-NMe₂

L12 ANSWER 7 OF 15 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 1995:938113 CAPLUS

DOCUMENT NUMBER: 123:332082

TITLE: Preparation of biotin derivative and method for

non-isotopic labeling of genes by biotin derivative

INVENTOR(S): Yamamoto, Isamu; Mukai, Tsunehiro

CODEN: JKXXAF

PATENT ASSIGNEE(S): Yamamoto Isamu, Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 5 pp.

DOCUMENT TYPE: Patent

LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

М

the

PATENT NO. KIND DATE APPLICATION NO. DATE

JP 07157497 A2 19950620 JP 1993-330034 19931201

OTHER SOURCE(S): MARPAT 123:332082

AB A carbodiimide-contg. biotin deriv. (I; R1 = C1-6 alkyl, cycloalkyl; R2 = C1-6 alkylene; R3, R4 = C1-3 alkyl; X- = C1-, Br-, or I-) is prepd. A non-isotopic labeling of a gene involves biotinylation of a DNA or RNA by reacting a DNA or RNA with a biotin deriv. having a carbodiimide group I. The biotin deriv. can be prepd. in relatively low cost, readily reacts with a DNA or RNA, and the reaction product is colored and can be distinguished from other non-labeled compds., DNA, or RNA. Thus, 260 mg biotin hydrazide was dissolved in 10 mL 0.5M NaHCO3, followed by adding a soln. of bromoacetic anhydride in dioxane at 0.degree., and after 15 min, the formed ppt. was filtered and recrystd. from H2O to give 227.4 mg biotin N-bromoacetylhydrazide. The latter compd. (0.76 g) and 0.31 g 1-ethyl-3-(3-dimethylaminopropyl)carbodiimide were added to 10 mL DMF and the formed ppt. was filtered, washed with Et2O, and dried in vacuo to

give 100% I [R1 = Et, R2 = (CH2)3, R3 = R4 = Me, X = Br] (II). A single strand

of DNA of M13mp18 (5 .mu.g) was dissolved in .apprx.5 .mu.L 0.1 M boric acid buffer (pH 8.0) and mixed with a soln. of the carbodiimide II (50 .mu.g/.mu.L) in the same buffer (5 .mu.L) and the mixt. was allowed to react at 37.degree. for 2 h. To the reaction mixt. was added 10 .mu.L 5

AcONH4 buffer and 60 .mu.L EtOH was added to ppt. biotinylated DNA, which was removed by filtration and dissolved in 10 .mu.L H2O. According to

measurement by UV absorption (260 nm), 4.5 .mu.g DNA was recovered. The recovered DNA was dild. to 1-128 pg/.mu.L and each soln. was spotted on a nitro cellulose filter and successively reacted with a streptoavidin-alkali phosphatase conjugate, NBT, and BCIP. The each spot

```
was detected at least 1 pg/.mu.L by blue coloration. II was also used
for
    non-isotopic labeling of DNA probes in the southern hybridization method.
     32897-26-0P, 1-Ethyl-3-(3-dimethylaminopropyl)urea
ΙT
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
        (intermediate for prepn. of carbodiimide-contg. biotin deriv. for
        non-isotopic labeling of DNA and RNA)
     32897-26-0 CAPLUS
RN
     Urea, N-[3-(dimethylamino)propyl]-N'-ethyl- (9CI) (CA INDEX NAME)
CN
     0
EtNH-C-NH-(CH<sub>2</sub>)<sub>3</sub>-NMe<sub>2</sub>
L12 ANSWER 8 OF 15 CAPLUS COPYRIGHT 2001 ACS
                         1995:785100 CAPLUS
ACCESSION NUMBER:
                         123:193056
DOCUMENT NUMBER:
                         Non-specific reaction suppressor for immunoassays
TITLE:
                         Ito, Michio; Sugawa, Satoshi; Yanagida, Atsushi
INVENTOR(S):
                         Mitsubishi Chemical Corp., Japan
PATENT ASSIGNEE(S):
                         Eur. Pat. Appl., 20 pp.
SOURCE:
                         CODEN: EPXXDW
                         Patent
DOCUMENT TYPE:
                         English
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                                           APPLICATION NO. DATE
                      KIND
                            DATE
     PATENT NO.
                                           _____
                                           EP 1995-101638
                                                            19950207
     EP 667529
                      A2
                            19950816
                            19960124
     EP 667529
                      A3
         R: DE, FR, GB, IT
                                           US 1994-194475
                                                            19940209
                            19960409
     US 5506151
                      Α
                                                            19950208
                                           CN 1995-102794
                            19951101
     CN 1111016
                       A
                                           JP 1995-22072
                                                             19950209
     JP 07253430
                            19951003
                      A2
                                                             19940209
                                        US 1994-194475
PRIORITY APPLN. INFO.:
                         MARPAT 123:193056
OTHER SOURCE(S):
     Disclosed is a non-specific reaction suppressor for immunoassays having
```

the formula: R1R2N(CHY)m(X)n(CHY)pR3, where R1, R2 = C1-5 alky1; X = -NHCONH-, -NHCSNH-, etc; Y = H, OH, or halogen; and R3 = NH2, NR1R2, cyclohexyl, or H; m = 0-5; p = 0-5; and n = 0 or 1. Also disclosed is a immunoassay uses latex particle-immobilized immunoreactant and nonspecific

reaction suppressor, e.g. 1-ethyl-3-(3-dimethyl-aminopropyl)urea, 1-cyclohexyl-3-3(2-morpholinoethyl)urea metho-p-toluenesulfone, dimethylamine, etc. In example, latex-immobilized digoxin, anti-digoxin antibody reagent compn., and EDU contg. 1-ethyl3-(3-dimethylaminopropyl)carbodiimide HCl were prepd. and tested.

32897-26-0, 1-Ethyl3-(3-dimethylaminopropyl)urea ΙT RL: MOA (Modifier or additive use); USES (Uses) (immunoassay uses latex particle-immobilized immunoreactant and nonspecific reaction suppressor)

32897-26-0 CAPLUS RN

09/350,193

CM 2

CRN 88-89-1 CMF C6 H3 N3 O7

L23 ANSWER 10 OF 10 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

1978:590538 CAPLUS

DOCUMENT NUMBER:

89:190538

TITLE:

Method for the photometric determination of

N-monosubstituted carbamates

AUTHOR(S):

Schoene, K.; Steinhanses, J.

CORPORATE SOURCE: SOURCE:

Inst. Aerobiol., Fraunhofer-Ges., Schmallenberg, Ger.

Fresenius' Z. Anal. Chem. (1978), 292(1), 29-33

CODEN: ZACFAU; ISSN: 0016-1152

DOCUMENT TYPE:

Journal

LANGUAGE:

German

AB N-monosubstituted carbamates were converted to urea derivs. by reaction with 1,3-diaminopropane in the presence of small amts. of NaOH. The urea derivs. were detd. spectrophotometrically at 577 nm by using the carbamide

reaction described by W. R. Fearon (1939). The detection limit for N-methylcarbamates is in the range of 20 nmols. In the case of N-methylcarbamates, N-methyl-N'-(3-aminopropyl)urea was found to be the intermediate urea deriv., which is formed in nearly quant. yield. The amidation reaction mechanism of the N-methylcarbamates was studied on N-methylurethane.

IT 68156-37-6P

RN 68156-37-6 CAPLUS

CN Urea, N-(3-aminopropyl)-N'-methyl- (9CI) (CA INDEX NAME)

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CA SUBSCRIBER PRICE

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=> d his (FILE 'HOME' ENTERED AT 10:38:39 ON 01 JUN 2001) FILE 'REGISTRY' ENTERED AT 10:38:44 ON 01 JUN 2001 STRUCTURE UPLOADED L1L2 50 S L1 L3 36270 S L1 FULL STRUCTURE UPLOADED 14060 S L4 FULL SUB=L3 L55399 S L5 AND 3/N L6 734 S ETHYL(L) DIMETHYL(L) AMINO(L) PROPYL(L) UREA L7 0 S UREA, "N-ETHYL-N'-(3-DIMETHYLAMINOPROPYL)-" L8 4 S ETHYL(L) DIMETHYLAMINOPROPYL(L) UREA L9 L103 S L9 AND 1/NC 1 S L10 AND 1/0 L11 FILE 'CAPLUS' ENTERED AT 10:52:57 ON 01 JUN 2001 15 S L11 L122 S L11/THU L13 FILE 'USPATFULL' ENTERED AT 10:55:19 ON 01 JUN 2001 L14 2 S L11 FILE 'MARPAT' ENTERED AT 10:56:49 ON 01 JUN 2001 0 S L11 L15 0 S L9 L16 FILE 'BEILSTEIN' ENTERED AT 10:57:15 ON 01 JUN 2001 L17 1 S L11 FILE 'REGISTRY' ENTERED AT 10:57:49 ON 01 JUN 2001 STRUCTURE UPLOADED L18 L19 11247 S L18 FULL SUB=L3 50 S L18 L20 0 S L18 CSS L21 8 S L18 CSS FULL L22 FILE 'CAPLUS' ENTERED AT 11:01:43 ON 01 JUN 2001 L23 10 S L22 => s 122/thu 10 L22 375285 THU/RL 0 L22/THU L24 (L22 (L) THU/RL) => file uspatful SINCE FILE COST IN U.S. DOLLARS TOTAL ENTRY SESSION

44.43

ENTRY

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SINCE FILE

598.50

TOTAL

SESSION

-15.88

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PASSWORD:

TERMINAL (ENTER 1, 2, 3, OR ?):2

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NEWS WWW CAS World Wide Web Site (general information)

Enter NEWS followed by the item number or name to see news on that specific topic.

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COST IN U.S. DOLLARS

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0.15 0.15

FULL ESTIMATED COST

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TSCA INFORMATION NOW CURRENT THROUGH January 11, 2001

Please note that search-term pricing does apply when

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Structure search limits have been increased. See HELP SLIMIT for details.

=>

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L1 STRUCTURE UPLOADED

=> s 11

SAMPLE SEARCH INITIATED 10:39:07 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 3038 TO ITERATE

32.9% PROCESSED 1000 ITERATIONS INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)

INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)

SEARCH TIME: 00.00.02

FULL FILE PROJECTIONS: ONLINE **COMPLETE**

BATCH **COMPLETE**

50 ANSWERS

PROJECTED ITERATIONS: 57457 TO 64063 PROJECTED ANSWERS: 33720 TO 38826

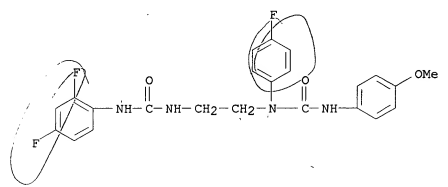
L2 50 SEA SSS SAM L1

=> d scan

L2 50 ANSWERS REGISTRY COPYRIGHT 2001 ACS

IN Urea, N-[2-[[[(2,4-difluorophenyl)amino]carbonyl]amino]ethyl]-N-(4fluorophenyl)-N'-(4-methoxyphenyl)- (9CI)

MF C23 H21 F3 N4 O3



HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):1

L2 50 ANSWERS REGISTRY COPYRIGHT 2001 ACS

IN Urea, N-[3-[[(3-methoxyphenyl)amino]carbonyl]amino]propyl]-N'-(4-methylphenyl)-N-(4-phenoxyphenyl)- (9CI)

MF C31 H32 N4 O4

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):1

L2 50 ANSWERS REGISTRY COPYRIGHT 2001 ACS

IN Propanamide,

3-[[(3-methoxyphenyl)amino]carbonyl](1-phenylethyl)amino]-N-(2-phenylethyl)-N-(phenylmethyl)- (9CI)

MF C34 H37 N3 O3

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):1

L2 50 ANSWERS REGISTRY COPYRIGHT 2001 ACS

IN Urea, N-[6-[(4-nitrophenyl)amino]hexyl]-N'-(phenylmethyl)- (9CI)

MF C20 H26 N4 O3

$$O = 0$$
 $O = 0$
 $O =$

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):2

L2 50 ANSWERS REGISTRY COPYRIGHT 2001 ACS

IN Urea, N'-(4-bromophenyl)-N-[(2-fluorophenyl)methyl]-N-[6-(2pyridinylamino)hexyl]- (9CI)

MF C25 H28 Br F N4 O

09/350,193

L2 50 ANSWERS REGISTRY COPYRIGHT 2001 ACS

MF C22 H17 C1 F N5 O6

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

=> d his

(FILE 'HOME' ENTERED AT 10:38:39 ON 01 JUN 2001)

FILE 'REGISTRY' ENTERED AT 10:38:44 ON 01 JUN 2001

L1 STRUCTURE UPLOADED

L2 50 S L1

=> s l1 full

FULL SEARCH INITIATED 10:40:07 FILE 'REGISTRY'
FULL SCREEN SEARCH COMPLETED - 60020 TO ITERATE

100.0% PROCESSED 60020 ITERATIONS

SEARCH TIME: 00.00.08

36270 ANSWERS

L3 36270 SEA SSS FUL L1

=>

Uploading 489.str

L4 STRUCTURE UPLOADED

=> s 14 sub=13 full FULL SUBSET SEARCH INITIATED 10:40:55 FILE 'REGISTRY' FULL SUBSET SCREEN SEARCH COMPLETED - 32305 TO ITERATE

100.0% PROCESSED 32305 ITERATIONS SEARCH TIME: 00.00.04

14060 ANSWERS

L5 14060 SEA SUB=L3 SSS FUL L4

=> d scan

L5 14060 ANSWERS REGISTRY COPYRIGHT 2001 ACS

IN INDEX NAME NOT YET ASSIGNED

MF C24 H30 Cl2 N4 O6 S

Absolute stereochemistry.

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

=> s 15 and 3/n 1884504 3/N L6 5399 L5 AND 3/N

=> d scan

L6 5399 ANSWERS REGISTRY COPYRIGHT 2001 ACS
IN Urea, N-[3-(dimethylamino)propyl]-N'-(2-phenoxyphenyl)- (9CI)
MF C18 H23 N3 O2

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):2

L6 5399 ANSWERS REGISTRY COPYRIGHT 2001 ACS
IN Heptanamide,
N-[2-[(4-fluorophenyl)[[(2-fluorophenyl)amino]carbonyl]amino]
 ethyl]- (9CI)
MF C22 H27 F2 N3 O2

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

=> s ethyl(1)dimethyl(1)amino(1)propyl(1)urea 4484821 ETHYL 11 ETHYLS 4484821 ETHYL (ETHYL OR ETHYLS) 2862325 DIMETHYL 3 DIMETHYLS 2862325 DIMETHYL (DIMETHYL OR DIMETHYLS) 3229260 AMINO 8155 AMINOS 3229260 AMINO (AMINO OR AMINOS) 1471974 PROPYL 4 PROPYLS 1471974 PROPYL (PROPYL OR PROPYLS)

09/350,193

162482 UREA

1 UREAS

162482 UREA

(UREA OR UREAS)

L7 734 ETHYL(L) DIMETHYL(L) AMINO(L) PROPYL(L) UREA

=> d scan

L7 734 ANSWERS REGISTRY COPYRIGHT 2001 ACS

Urea, N'-(4-chlorophenyl)-N-[3-[[[(1,1-dimethylethyl)amino]carbonyl]a
mino]propyl]-N-(4-fluorophenyl)- (9CI)

MF C21 H26 C1 F N4 O2

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):5

L7 734 ANSWERS REGISTRY COPYRIGHT 2001 ACS

Urea, N'-(4-chlorophenyl)-N-[1-[3,4-dihydro-3-(3-methylphenyl)-4-oxo2-quinazolinyl]propyl]-N-[2-(dimethylamino)ethyl]- (9CI)

MF C29 H32 C1 N5 O2

$$\begin{array}{c|c} & \text{CH}_2\text{-}\text{CH}_2\text{-}\text{NMe}_2\\ & \text{Et} & \text{O}\\ & \text{CH}-\text{N}-\text{C}-\text{NH} \end{array}$$

L7 734 ANSWERS REGISTRY COPYRIGHT 2001 ACS

IN Urea, N-[1-(1,4-dihydro-4-oxo-2-quinazolinyl)propyl]-N-[2(dimethylamino)ethyl]-N'-(2-fluorophenyl)- (9CI)

MF C22 H26 F N5 O2

09/350,193

$$\begin{array}{c|c} H & \text{Et} & \text{CH}_2\text{--}\text{CH}_2\text{--}\text{NMe}_2 \\ \hline & | & | & | \\ N & \text{CH}\text{--}\text{N}\text{---}\text{R} \\ \hline & O \end{array}$$

L7 734 ANSWERS REGISTRY COPYRIGHT 2001 ACS

L7 734 ANSWERS REGISTRY COPYRIGHT 2001 ACS

Relative stereochemistry.

●2 HCl

L7 734 ANSWERS REGISTRY COPYRIGHT 2001 ACS
IN Urea, 1-[2-(dimethylamino)ethyl]-1-isopentyl-3,3-diisopropyl(7CI)

MF C16 H35 N3 O

$$\begin{tabular}{l} \tt O \\ || \\ \tt C-N(Pr-i)_2 \\ | \\ \tt Me_2N-CH_2-CH_2-N-CH_2-CH_2-CHMe_2 \\ \end{tabular}$$

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):5

L7 734 ANSWERS REGISTRY COPYRIGHT 2001 ACS

IN Urea, N'-[4-[3-[(1,1-dimethylethyl)amino]-2-hydroxypropoxy]-3-(1oxopropyl)phenyl]-N,N-dimethyl-, (2E)-2-butenedioate (1:1) (salt) (9CI)
MF C19 H31 N3 O4 . C4 H4 O4

CM 1

$$\begin{array}{c|c} \text{O} \\ \text{NH-C-NMe}_2 \\ \\ \text{OH} \\ \text{C-Et} \\ \\ \text{t-BuNH-CH}_2\text{-CH-CH}_2\text{-O} \\ \\ \text{O} \end{array}$$

CM 2

Double bond geometry as shown.

L7 734 ANSWERS REGISTRY COPYRIGHT 2001 ACS

Urea, N'-[2,6-bis(1-methylethyl)phenyl]-N-[2-(dimethylamino)ethyl]-N[1-[3-(4-ethylphenyl)-3,4-dihydro-4-oxo-2-quinazolinyl]propyl]- (9CI)
MF C36 H47 N5 O2

L7 734 ANSWERS REGISTRY COPYRIGHT 2001 ACS

Urea, N-[1-[3-(4-chlorophenyl)-3,4-dihydro-4-oxo-2quinazolinyl]propyl]-N-[2-(dimethylamino)ethyl]-N'-(3-methylphenyl)(9CI)

MF C29 H32 C1 N5 O2

L7 734 ANSWERS REGISTRY COPYRIGHT 2001 ACS

IN Urea, N'-cyclohexyl-N-[1-(3,4-dihydro-4-oxo-3-phenyl-2quinazolinyl)propyl]-N-[2-(dimethylamino)ethyl]- (9CI)
MF C28 H37 N5 O2

L7 734 ANSWERS REGISTRY COPYRIGHT 2001 ACS

Urea, N-(2-cyclohexylethyl)-N'-[3-(dimethylamino)propyl]-N-(2hydroxyethyl)- (9CI)

MF C16 H33 N3 O2

$$\begin{array}{c} \text{CH}_2-\text{CH}_2-\text{OH} \\ | \\ \text{CH}_2-\text{CH}_2-\text{N}-\text{C}-\text{NH}-\text{(CH}_2)}_3-\text{NMe}_2 \\ | \\ | \\ \text{O} \end{array}$$

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

=> s urea, N-ethyl-N'-(3-dimethylaminopropyl)MISMATCHED QUOTE 'N-ETHYL-N'-'
Quotation marks (or apostrophes) must be used in pairs,

```
one before and one after the expression you are setting
off or masking.
=> s urea, "N-ethyl-N'-(3-dimethylaminopropyl)-"
         162482 UREA
              1 UREAS
         162482 UREA
                  (UREA OR UREAS)
       3623758 "N"
       4484821 "ETHYL"
             11 "ETHYLS"
       4484821 "ETHYL"
                  ("ETHYL" OR "ETHYLS")
        529941 "N'"
       9790704 "3"
           1914 "DIMETHYLAMINOPROPYL"
L8
              O UREA, "N-ETHYL-N'-(3-DIMETHYLAMINOPROPYL)-"
                  (UREA(W) "N"(W) "ETHYL"(W) "N'"(W) "3"(W) "DIMETHYLAMINOPROPYL")
=> s ethyl(l)dimethylaminopropyl(l)urea
       4484821 ETHYL
             11 ETHYLS
       4484821 ETHYL
                  (ETHYL OR ETHYLS)
           1914 DIMETHYLAMINOPROPYL
        162482 UREA
              1 UREAS
        162482 UREA
                  (UREA OR UREAS)
L9
              4 ETHYL (L) DIMETHYLAMINOPROPYL (L) UREA
=> d scan
L9
     4 ANSWERS
                  REGISTRY COPYRIGHT 2001 ACS
IN
     Sulfamic acid, cyclohexyl-, compd. with N-[2-[[3-
     (dimethylamino)propyl]thio]phenyl]-N'-ethylurea (1:1) (9CI)
     C14 H23 N3 O S . C6 H13 N O3 S
MF
     CM
          1
       S-(CH<sub>2</sub>)<sub>3</sub>-NMe<sub>2</sub>
           -C-NHEt
           0
```

CM 2

09/350,193

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

=> s 19 and 1/nc 27444421 1/NC L10 3 L9 AND 1/NC

=> d scan

L10 3 ANSWERS REGISTRY COPYRIGHT 2001 ACS
IN 2-Propenoic acid, 2-methyl-,
2-[[[[3-(dimethylamino)propyl]amino]carbonyl]
 amino]ethyl ester (9CI)
MF C12 H23 N3 O3

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):2

L10 3 ANSWERS REGISTRY COPYRIGHT 2001 ACS
IN Urea, N-[3-(dimethylamino)propyl]-N'-ethyl- (9CI)
MF C8 H19 N3 O
CI COM

L10 3 ANSWERS REGISTRY COPYRIGHT 2001 ACS
IN Thiourea, N-[3-(dimethylamino)propyl]-N'-ethyl- (9CI)
MF C8 H19 N3 S

ALL ANSWERS HAVE BEEN SCANNED

```
=> s Urea, N-[3-(dimethylamino)propyl]-N'-ethyl-/cn
MISMATCHED QUOTE 'PROPYL] -N'-ETHYL-/CN'
Quotation marks (or apostrophes) must be used in pairs,
one before and one after the expression you are setting
off or masking.
=> s 110 and C8 H19 N3 O
           1563 C8
             67 H19
          57156 N3
       1833420 O
              0 C8 H19 N3 O
                   (C8(W)H19(W)N3(W)O)
L11
              0 L10 AND C8 H19 N3 O
=> s C8 H19 N3 O
           1563 C8
             67 H19
          57156 N3
       1833420 O
              0 C8 H19 N3 O
L12
                  (C8(W)H19(W)N3(W)O)
=> del 111- y
=> d his
     (FILE 'HOME' ENTERED AT 10:38:39 ON 01 JUN 2001)
     FILE 'REGISTRY' ENTERED AT 10:38:44 ON 01 JUN 2001
L1
                 STRUCTURE UPLOADED
L2
              50 S L1
L3
           36270 S L1 FULL
                 STRUCTURE UPLOADED
L4
L5
          14060 S L4 FULL SUB=L3
L6
            5399 S L5 AND 3/N
             734 S ETHYL(L)DIMETHYL(L)AMINO(L)PROPYL(L)UREA
L7
               0 S UREA, "N-ETHYL-N'-(3-DIMETHYLAMINOPROPYL)-"
L8
               4 S ETHYL(L)DIMETHYLAMINOPROPYL(L)UREA
L9
               3 S L9 AND 1/NC
L10
\Rightarrow s 110 and 1/o
       2846307 1/0
L11
              1 L10 AND 1/0
=> d scan
                  REGISTRY COPYRIGHT 2001 ACS
L11 1 ANSWERS
IN
     Urea, N-[3-(dimethylamino)propyl]-N'-ethyl- (9CI)
MF
     C8 H19 N3 O
CI
     COM
      0
EtNH-C-NH-(CH<sub>2</sub>)<sub>3</sub>-NMe<sub>2</sub>
```

O || EtNH-C-NH-(CH₂)₃-NMe₂

ALL ANSWERS HAVE BEEN SCANNED

=> file caplus
COST IN U.S. DOLLARS

FULL ESTIMATED COST

SINCE FILE TOTAL ENTRY SESSION 267.19 267.34

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=> d his

L1

L2

(FILE 'HOME' ENTERED AT 10:38:39 ON 01 JUN 2001)

FILE 'REGISTRY' ENTERED AT 10:38:44 ON 01 JUN 2001 STRUCTURE UPLOADED 50 S L1

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L3
          36270 S L1 FULL
L4
                STRUCTURE UPLOADED
L5
          14060 S L4 FULL SUB=L3
L6
           5399 S L5 AND 3/N
L7
            734 S ETHYL(L) DIMETHYL(L) AMINO(L) PROPYL(L) UREA
L8
              0 S UREA, "N-ETHYL-N'-(3-DIMETHYLAMINOPROPYL)-"
              4 S ETHYL(L)DIMETHYLAMINOPROPYL(L)UREA
L9
              3 S L9 AND 1/NC
L10
L11
              1 S L10 AND 1/0
     FILE 'CAPLUS' ENTERED AT 10:52:57 ON 01 JUN 2001
=> s 111
            15 L11
L12
=> d ibib ab hitstr 1-15
L12 ANSWER 1 OF 15 CAPLUS COPYRIGHT 2001 ACS
ACCESSION NUMBER:
                         2001:338479 CAPLUS
                         Preparation of amides and ureas as activators of
TITLE:
                         soluble guanylate cyclase
INVENTOR(S):
                         Selwood, David; Glen, Robert; Reynolds, Karen;
                         Wishart, Grant
PATENT ASSIGNEE(S):
                         University College London, UK
SOURCE:
                         PCT Int. Appl., 101 pp.
                         CODEN: PIXXD2
DOCUMENT TYPE:
                         Patent
LANGUAGE:
                         English
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
     PATENT NO.
                    KIND DATE
                                          APPLICATION NO. DATE
     ----- ---- ----
                           _____
                                          -----
     WO 2001032604
                     A1 20010510
                                         WO 2000-GB4249 20001106
        W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
             CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR,
             HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT,
             LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU,
             SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN,
             YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
             DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,
             BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
PRIORITY APPLN. INFO.:
                                        GB 1999-26286
                                                        A 19991105
                                        US 2000-201382
                                                        P 20000502
AΒ
     The title compds. R4PZNR1R2 [I; R1, R2 = alkyl; R1R2 together form
     alkylene; Z = alkylene; P = a direct bond, X, Y, W, XY, YW, XYW (wherein
W
     = O, S, NR3; R3 = H, alkyl; Y = UV; V = a direct bond, alkylene; U = CS,
    CO, SO2, C(:NR); R = H, OH, alkyl; X = O, NR6; R6 = H, alkyl, alkenyl,
    etc.); R4 = alkyl, alkenyl, alkynyl, etc.], useful in the activation of
    sol. guanylate cyclase, were prepd. E.g., synthesis of the urea II,
    starting with 4-bromoaniline and 1-(3-aminopropyl)pyrrolidine, was given.
    Biol. data for compds. I (e.g., IC50 for inhibition of platelet
    aggregation) were presented.
ΙT
    32897-26-0P
```

RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of amides and ureas as activators of sol. guanylate cyclase)

RN32897-26-0 CAPLUS

CN Urea, N-[3-(dimethylamino)propyl]-N'-ethyl- (9CI) (CA INDEX NAME)

0 EtNH-C-NH-(CH₂)₃-NMe₂

REFERENCE COUNT: 24

REFERENCE(S):

1986.

(8) Catanese, B; BOLLETTINO CHIMICO FARMACEUTICO

V125(7), P228 CAPLUS

(9) Farbenfabriken Bayer Aq; DE 890958 C CAPLUS

(10) Glen, R; WO 0027394 A 2000 CAPLUS

(12) Hoechst Marion Roussel de Gmbh; EP 0908456 A

1999

CAPLUS

(13) Hoechst Marion Roussel de Gmbh; DE 19756388 A

1999 CAPLUS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L12 ANSWER 2 OF 15 CAPLUS COPYRIGHT 2001 ACS 2000:852726 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER:

134:243293

TITLE:

A cyclohexane-1,2-diyldinitrilotetraacetate tetrahydroxamate derivative for actinide

complexation:

synthesis and complexation studies

AUTHOR(S):

Santos, M. Amelia; Rodrigues, Estela; Gaspar,

Margarida

CORPORATE SOURCE:

Centro de Quimica Estrutural, Complexo I, Instituto

Superior Tecnico, Lisbon, 1049-001, Port.

SOURCE:

Dalton (2000), (23), 4398-4402

CODEN: DALTFG

PUBLISHER:

Royal Society of Chemistry

DOCUMENT TYPE:

Journal

LANGUAGE:

English

A new tetrahydroxamate ligand has been synthesized and its chelating properties studied, in aq. solns., with thorium(IV) and iron(III) as analogs of the actinides plutonium(IV) and (to some extent) americium(III). The architecture of this ligand is based on that of the cyclohexane-1,2-diyldinitrilotetraacetate complexon with hydroxamate instead of carboxylate groups. It has proven to form quite stable and water sol. complexes with these metal ions, up to pH 9. Besides the 1:1 (M:L) monomeric species formed under acidic conditions, the corresponding (2:2) dimeric complexes may also be admitted under physiol. conditions. According to the magnetic properties and modeling calcns., the iron(III) dimer species should have some magnetic interaction between the metallic centers.

IT 32897-26-0

RL: RCT (Reactant)

```
(starting material in prepn. of cyclohexane-1,2-diyldinitrilotetra(N-
          methylacetohydroxamic acid))
RN
      32897-26-0 CAPLUS
      Urea, N-[3-(dimethylamino)propyl]-N'-ethyl- (9CI) (CA INDEX NAME)
CN
       0
EtNH-C-NH-(CH_2)_3-NMe_2
REFERENCE COUNT:
                               37
                               (4) Bouby, M; J Alloys Comp 1998, V271-273, P206
REFERENCE(S):
                                    CAPLUS
                               (5) Carrano, C; J Am Chem Soc 1979, V101, P599 CAPLUS
                               (6) Dasaradhi, L; J Chem Soc, Perkin Trans 2 1997,
                                    P1187 CAPLUS
                               (8) Esteves, M; J Chem Soc, Dalton Trans 1995, P2565
                                    CAPLUS
                               (9) Evans, D; J Chem Soc 1959, P2003 CAPLUS
                               ALL CITATIONS AVAILABLE IN THE RE FORMAT
L12 ANSWER 3 OF 15 CAPLUS COPYRIGHT 2001 ACS
                            2000:725451 CAPLUS
ACCESSION NUMBER:
DOCUMENT NUMBER:
                               133:286497
TITLE:
                               Immunomodulatory compositions and methods of use
                               thereof
INVENTOR(S):
                               Onderdonk, Andrew B.; Tzianabos, Arthur O.; Miller,
                               Robert J.; Calias, Pericles
PATENT ASSIGNEE(S):
                               Genzyme Corporations, USA
                               PCT Int. Appl., 62 pp.
SOURCE:
                               CODEN: PIXXD2
DOCUMENT TYPE:
                               Patent
LANGUAGE:
                               English
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                                                     APPLICATION NO.
      PATENT NO.
                           KIND
                                  DATE
                                                                          DATE
                           ____
                                  _____
                                                     ₩<mark>Ò</mark>,2000-US9087
      WO 2000059490
                           A2
                                  20001012
                                                                          20000406
      WO 2000059490
                           A3
                                  20010215
               AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR,
                CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU,
                LV, MA, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZW,
          SG, SI, SK, SL, TJ, TM, IK, LI, IZ, GA, GG, GZ, MA, AZ, BY, KG, KZ, MD, RW. TJ, TM

RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, TE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG

APPLN. INFO::

US 1999-128177 P 19990406
PRIORITY APPLN. INFO.:
OTHER SOURCE(S):
                              MARPAT 133:286497
      The invention relates to immunomodulatory compns. and related methods.
AB
      The immunomodulatory compnstare useful for the prevention of sepsis and
      the treatment and prevention of diseases assocd. with inflammation and/or
             CM-cellulose/N-ethyl-N'\(3-dimethylaminopropyl)urea formulations
are
```

```
described.
IT
     32897-26-0
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (immunomodulatory compns.)
RN
     32897-26-0 CAPLUS
CN
     Urea, N-[3-(dimethylamino)propyl]-N'-ethyl- (9CI) (CA INDEX NAME)
      0
EtNH-C-NH-(CH<sub>2</sub>)<sub>3</sub>-NMe<sub>2</sub>
L12 ANSWER 4 OF 15 CAPLUS COPYRIGHT 2001 ACS
ACCESSION NUMBER:
                         2000:368150 CAPLUS
DOCUMENT NUMBER:
                         133:12765
                         Preventives and/or remedies for central nervous
TITLE:
system
                         diseases containing compounds having TXA2 receptor
                         antagonism and/or TXA2/synthase inhibitory effect
                         Yagami, Tatsuro; Honma, Tsunetoshi; Katsuura, Goro
INVENTOR(S):
                         Shionogi & Co., Ltd. / Japan
PATENT ASSIGNEE(S):
                         PCT Int. Appl., 171 pp.
SOURCE:
                         CODEN: PIXXD2
DOCUMENT TYPE:
                         Patent
LANGUAGE:
                         Japanese
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
     PATENT NO.
                      KIND
                            DATE
                                            APPLICATION NO.
                                                             DATE
     WO 2000030683
                            20000602
                      A1
                                           WO 1999-JP6317
                                                             19991112
         W: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU,
             CZ, DE, DK, DM, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL,
             IN, IS, JP, KE, KG, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD,
             MG, MK, MN, MW, MX, NO,/NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK,
             SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ,
             BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE,
             DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF,
             CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
PRIORITY APPLN. INFO.:
                                        JP 1998-329862
                                                          A 19981119
     Compds. having TXA2 antagonism and/or a TXA2 synthase inhibitory effect,
     prodrugs thereof, pharmaceutically acceptable salts of the same or
     hydrates of the same, which show effects of inhibiting nerve cell
     denaturation caused by amyloid .beta. protein and nerve cell death caused
    by axonotmesis, are useful as preventives and/or remedies for central
     nervous system diseases, preventives and/or remedies for nerve
     degeneration diseases, nerve cell denaturation inhibitors, amyloid .beta.
     protein-induced nerve cell denaturation inhibitors, nerve cell death
     inhibitors, axonotmesis-induced nerve cell death inhibitors and, in
    particular, preventives and/or remedies for dementia of Alzheimer type.
IT
     32897-26-0
     RL: RCT (Reactant)
        (preventives and/or remedies for central nervous system diseases
contq.
```

```
compds. having TXA2 receptor antagonism and/or TXA2 synthase
inhibitory
        effect)
RN
     32897-26-0 CAPLUS
     Urea, N-[3-(dimethylamino)propyl]-N'-ethyl- (9CI) (CA INDEX NAME)
CN
      0
EtNH-C-NH-(CH<sub>2</sub>)<sub>3</sub>-NMe<sub>2</sub>
REFERENCE COUNT:
                          (1) Arimura, A; Br J Pharmacol 1998, ¥124, P795
REFERENCE(S):
CAPLUS
                          (2) Dickinson, R; Bioorg Med Chem Lett 1996, V6(14),
                              P1691 CAPLUS
                          (3) Hall, S; Med Res Rev 1991, V11(5), P503 CAPLUS
                          (6) Shionogi & Co Ltd; GB 2184118 A CAPLUS
                          (7) Shionogi & Co Ltd; US 4960909 A CAPLUS
                          ALL CITATIONS AVAILABLE IN THE RE FORMAT
L12 ANSWER 5 OF 15 CAPLUS COPYRIGHT 2001 ACS
ACCESSION NUMBER:
                          2000:277959 CAPLUS
DOCUMENT NUMBER:
                          132:321662
TITLE:
                          Preparation of aromatic amine derivatives and agents
                          containing the same
INVENTOR(S):
                         Oi, Satoru; Suzuķi, Nobuhiro; Aso, Kazuyoshi; Banno,
                         Yoshihiro
PATENT ASSIGNEE(S):
                         Takeda Chemical Industries, Ltd., Japan
                         PCT Int. Appl/, 309 pp.
SOURCE:
                         CODEN: PIXXD2
DOCUMENT TYPE:
                         Patent
LANGUAGE:
                          Japanese
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
     PATENT NO.
                      KIND
                            DATÉ
                                            APPLICATION NO.
                                                             DATE
     WO 2000023420
                       A1
                             20000427
                                            WO 1999-JP5755
                                                              19991019
         W: AE, AL, AM, AU, AZ, BA, BB, BG, BR, BY, CA, CN, CR, CU, CZ, DM,
             EE, GD, GE, HR, HU, ID, IL, IN, IS, JP, KG, KR, KZ, LC, LK, LR,
             LT, LV, MA, MD, MG, MK, MN, MX, NO, NZ, PL, RO, RU, SG, SI, SK,
             SL, TJ, TM, TR, TT, TZ, UA, US, UZ, VN, YU, ZA, AM, AZ, BY, KG,
             KZ, MD, RU, ŤJ, TM
         RW: GH, GM, KE,/LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE,
             DK, ES, FI/FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF,
             CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
     AU 9961246
                      A1
                           20000508
                                            AU 1999-61246
                                                             19991019
     JP 2000191615
                       A2
                            20000711
                                            JP 1999-297129
                                                             19991019
PRIORITY APPLN. INFO' .:
                                         JP 1998-298940 A 19981020
                                         WO 1999-JP5755
                                                          W
                                                             19991019
OTHER SOURCE(S):
                        MARPAT 132:321662
    Title compds / [I; wherein A is an optionally substituted arom. ring; B is
     an optionally substituted cyclic hydrocarbon oxy group; Z is an
optionally
```

Urea, N-[3-(dimethylamino)propyl]-N'-ethyl- (9CI) (CA INDEX NAME) CN

0 EtNH-C-NH-(CH2)3-NMe2

L12 ANSWER 9 OF 15 CAPLUS COPYRIGHT 2001 ACS 1995:294129 CAPLUS

ACCESSION NUMBER:

122:290591 DOCUMENT NUMBER:

Preparation of carbodiimide-containing biotin TITLE:

derivatives as reagents for detecting point mutation

of gene and diagnosis of hereditary disease

Yamamoto, Isamu; Mukai, Tsunehiro INVENTOR(S):

Yamamoto Isamu, Japan PATENT ASSIGNEE(S):

Jpn. Kokai Tokkyo Koho, 6 pp. SOURCE:

CODEN: JKXXAF

Patent DOCUMENT TYPE: Japanese LANGUAGE:

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

APPLICATION NO. DATE KIND DATE PATENT NO. ----_____ _____ ___ A2 19940927 19930315 JP 1993-80196 JP 06271581

MARPAT 122:290591 OTHER SOURCE(S):

The title biotin derivs. (I; R1 = C1-6 alkyl, cycloalkyl; R2 = C1-6 alkylene; R3, R4 = C1-3 alkyl; X = halogen ion, suitable for chem. modification of genes, are prepd. The presence and position of point mutation in a gene is detd. by (1) mixing for hybridization each complimentary single strand of a normal gene and its corresponding gene assuming the presence of point mutations, (2) reacting the above biotin deriv. I, (3) adsorbing the biotin deriv.-bonded DNA to a agarose column contg. avidin or its analog, (3) eluting the column with a soln. of biotin, and (5) detg. the base sequence of the isolated DNA fragment. Diagnosis of a hereditary disease involves (1) mixing for hybridization each complimentary single strand of a normal gene and its corresponding gene assuming the presence of point mutation, (2) reacting the above biotin deriv. I, and (3) detecting the biotin deriv.-bonded DNA by luminescence or fluorescence using avidin or its analog, which confirms the presence of gene point mutations. Both complimentary single strands of a normal gene and its corresponding gene assuming the presence of

mutation are obtained by cutting genes with a restriction enzyme. The avidin deriv. is a streptoavidin-alkali phosphatase conjugate. These carbodiimide-contg. biotin derivs. I react with guanine (G) or thymine

(T) of a double stranded DNA having G-T or T-G mismatching. Thus, 260 mg biotin hydrazide was dissolved in 0.5 M NaHCO3 followed by adding a soln. of 520 mg bromoacetic anhydride in dioxane at 0.degree., filtering off

the pptd. crystals after 15 min, and recrystn. from H2O to give 227.4 mg N-biotinyl-N'-bromoacetylhydrazine which was stirred with 1-cyclohexyl-3-(3-dimethylaminopropyl)carbodiimide in DMF to give 97%

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09/350,193
     title compd. I [R1 = cyclohexyl, R2 = (CH2)3, R3 = R4 = Me, X- = Br-]
     (II). Aldolase genes were cut out from both plasmid pHAA47 contg. normal
     A-type aldolase gene and plasmid pHAdA526 contg. A-type aldolase gene
from
     a hemolytic anemia patent but lacking erythrocyte aldolase activity by
     restriction enzyme Xab and HindIII, resp., sepd. by a agarose
     electrophoresis, and each digested by restriction enzyme Rsal into 3 DNA.
     Both digested genes were heated in a hybridization buffer at 100.degree.
     for 10 min and left to stand at 42.degree. overnight followed by
adjusting
     the pH to 8.5 and reacting with II at 30.degree. for 30 min. DNA's were
     sepd. by pptn. with EtOH, dissolved in H2O, and passed to a avidin
     column followed by eluting the column with 1 mM aq. biotin to sep.
     II-bonded DNA. As expected, the 411 bp fragment was recovered and
     confirmed to contain a mutation with the 386th adenine replaced with
     guanine in the patient lacking aldolase activity.
     32897-26-0P, 1-Ethyl-3-(3-dimethylaminopropyl)urea
IT
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
        (intermediate for prepn. of carbodiimide-contg. biotin derivs. as
        reagents for detecting gene point mutation and diagnosis of hereditary
        disease)
     32897-26-0 CAPLUS
RN
     Urea, N-[3-(dimethylamino)propyl]-N'-ethyl- (9CI) (CA INDEX NAME)
CN
EtNH-C-NH-(CH<sub>2</sub>)<sub>3</sub>-NMe<sub>2</sub>
```

L12 ANSWER 10 OF 15 CAPLUS COPYRIGHT 2001 ACS

1992:489833 CAPLUS ACCESSION NUMBER:

117:89833 DOCUMENT NUMBER:

Preparation of water-soluble 1-ethyl-3-(3-TITLE:

dimethylaminopropyl) carbodiimide

Yoneyama, Takahiro; Odagiri, Masaki; Imanari, Makoto INVENTOR(S): Keishitsu Ryubun Shinyoto Kaihatsu Gijutsu Kankyu PATENT ASSIGNEE(S):

Kumiai, Japan

Jpn. Kokai Tokkyo Koho, 3 pp. SOURCE:

CODEN: JKXXAF

DOCUMENT TYPE:

Patent Japanese LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 04077464	A2	19920311	JP 1990-189414	19900719
US 5208378	Α	19930504	US 1991-732123	19910718
PRIORITY APPLN. INFO.	:		JP 1990-189414	19900719
OTHER SOURCE(S):	CA	SREACT 117:	89833	

The title compd. (I) is prepd. by addn. reaction of EtNCS and N, N-dimethyl-1, 3-propanediamine (II) in arom. hydrocarbon, then treatment of the obtained thiourea deriv. with dehydrosulfurization agents without

isolation from the reaction mixt. A soln. of EtNCS in PhMe was teated dropwise with a soln. of II in PhMe under ice cooling over 2 h, stirred at room temp. for 2 h, then treated with Pb304 for 3 h under reflux to give 64% I. IT 32897-26-0P RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn. and dehydrosulfurization of) 32897-26-0 CAPLUS RNUrea, N-[3-(dimethylamino)propyl]-N'-ethyl- (9CI) (CA INDEX NAME) CN 0 EtNH-C-NH-(CH2)3-NMe2 L12 ANSWER 11 OF 15 CAPLUS COPYRIGHT 2001 ACS 1988:163694 CAPLUS ACCESSION NUMBER: 108:163694 DOCUMENT NUMBER: Isolation and purification of proteolytic enzymes on TITLE: organo-silica supports with immobilized gramicidin S Ignatchenko, A. P.; Bogomaz, V. I.; Tugai, V. A.; AUTHOR(S): Chuiko, A. A. A. V. Palladin Inst. Biochem., Kiev, USSR CORPORATE SOURCE: Ukr. Biokhim. Zh. (1987), 59(6), 28-33 SOURCE: CODEN: UBZHD4; ISSN: 0201-8470 Journal DOCUMENT TYPE: Russian LANGUAGE: Biospecific sorbents for affinity chromatog. of proteolytic enzymes were synthesized by attaching the cyclopeptide antibiotic gramicidin S to organo-silica supports. Gramicidin S was attached to the organo-silica supports using glutaric aldehyde, p-benzoquinone, sol. and insol. carbodiimides. The sorbents prepd. by these methods were successfully applied for the purifn. of the crude pepsin from horse gastric juice and proteolytic complex produced by Acremonium chrysogenum. 32897-26-0 ΙT RL: RCT (Reactant) (crosslinking by, of gramicidin S to organo-silica supports, for proteinase purifn.) 32897-26-0 CAPLUS RN Urea, N-[3-(dimethylamino)propyl]-N'-ethyl- (9CI) (CA INDEX NAME) CN

O || EtNH-C-NH-(CH₂)₃-NMe₂

L12 ANSWER 12 OF 15 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

1987:9247 CAPLUS

DOCUMENT NUMBER:

106:9247

TITLE:

Analytical, toxicological and immunological consequences of the use of N-ethyl-N'-(3-

dimethylaminopropyl)carbodiimide as coupling reagent

for the preparation of meningococcal group C

polysaccharide-tetanus toxoid conjugate as vaccine

for

human use

AUTHOR(S):

Beuvery, E. C.; Speijers, G. J. A.; Lutz, B. I. G.; Freudenthal, D.; Kanhai, V.; Haagmans, B.; Derks, H.

J. G. M.

CORPORATE SOURCE:

Rijksinst. Volksgezond. Milieuhyg., Bilthoven, 3720,

Neth.

SOURCE:

Dev. Biol. Stand. (1986), 63 (Use Stand. Chem. Defined

Antigens), 117-28

CODEN: DVBSA3; ISSN: 0301-5149

DOCUMENT TYPE:

Journal

LANGUAGE:

English

AB For the prepn. of meningococcal group C polysaccharide-tetanus toxoid conjugate the reactive reagent

N-ethyl-N'-(dimethylaminopropyl)carbodiimid

e is used. The application of this reagent results in a no. of stable linkages (viz. "peptide" linkages between the polysaccharide and tetanus toxoid, intrachain ester linkages in the polysaccharide component and binding of the N-acylurea deriv. of the reagent) and less stable ones (viz. anhydride linkages). As a consequence of the reaction, the reagent is converted to a nonreactive urea deriv. The toxic properties of the reagent and of the converted reagent were studied. These properties do not contraindicate the use of the coupling reagent for the prepn. of vaccines for human use. In addn. anal. methods were developed for the quant. evaluation of the coupling reagent, the reaction products and for the N-acylurea deriv. of the reagent and of the residual reactivity of conjugates for primary aminogroups. Although no test was performed for the assay of ester linkages in the polysaccharide component of the conjugate, evidence is presented that such linkages may be present. results of the test for residual reactivity indicated a spontaneous rearrangement of linkages after the prepn. of the conjugate. In addn.

the

effect of the ratio of coupling reagent-to-polysaccharide and tetanus toxoid on antigenic and immunogenic activities of the conjugate was studied. An increase of the ratio resulted in a decrease of the

antigenic

activity of the polysaccharide component but in an increase of its immunogenic activity as to the induction of IgG antibodies to the polysaccharide. The immunogenic activity of the polysaccharide component correlated rather well with the antigenic activity measured in heterologous enzyme-linked immunosorbent assay using antibodies to both components.

IT 32897-26-0P

RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation) (prepn. and toxicity of)

RN 32897-26-0 CAPLUS

CN Urea, N-[3-(dimethylamino)propyl]-N'-ethyl- (9CI) (CA INDEX NAME)

O || EtNH-C-NH-(CH₂)₃-NMe₂ L12 ANSWER 13 OF 15 CAPLUS COPYRIGHT 2001 ACS 1984:34771 CAPLUS ACCESSION NUMBER: DOCUMENT NUMBER: 100:34771 Synthesis of phosphoramidates of mono- and TITLE: oligonucleotides in aqueous media Gottikh, M. B.; Ivanovskaya, M. G.; Shabarova, Z. A. AUTHOR(S): Chem. Dep., M. V. Lomonosov Moscow State Univ., CORPORATE SOURCE: Moscow, USSR Bioorg. Khim. (1983), 9(8), 1063-7 SOURCE: CODEN: BIKHD7 DOCUMENT TYPE: Journal LANGUAGE: Russian Phosphoramidates of mono- and oligonucleotides were prepd. in 85-100% yields in aq. media by condensation of nucleotide component with any primary or secondary amine in the presence of EtC:N:C(CH2)3NMe2 (I) at a pH of 1 unit less than pKa value of the reacting amine, 0.5-4 h for amines with pKa < 8 in 4-20 h for amines with pKa > 8. Thus, condensation of 20 mmol pdT with 3 mmol PhNH2 at pH 3.5 for 5 min in the presence of 0.5 mol I gave 100% of the corresponding phosphoramidate. IT 32897-26-0 RL: RCT (Reactant) (condensation of mono- and oligonucleotides with primary and secondary amines in presence of) 32897-26-0 CAPLUS RN Urea, N-[3-(dimethylamino)propyl]-N'-ethyl- (9CI) (CA INDEX NAME) CN 0 EtNH-C-NH-(CH2)3-NMe2 L12 ANSWER 14 OF 15 CAPLUS COPYRIGHT 2001 ACS ACCESSION NUMBER: 1983:71187 CAPLUS DOCUMENT NUMBER: 98:71187 Direct spectrophotometric observation of an TITLE: O-acvlisourea intermediate: concerted general acid catalysis in the reaction of acetate ion with a water-soluble carbodiimide Ibrahim, Ibrahim T.; Williams, Andrew AUTHOR(S): Chem. Lab., Univ. Kent, Canterbury, CT2 7NZ, UK CORPORATE SOURCE: J. Chem. Soc., Perkin Trans. 2 (1982), (11), 1459-66 SOURCE: CODEN: JCPKBH; ISSN: 0300-9580 DOCUMENT TYPE: Journal LANGUAGE: English Rate consts. for the formation and decompn. of intermediate O-acylisoureas from carbodiimide and carboxylic acids were measured in aq. media. O-acetylisourea from AcO- and N-ethyl-N'-[3-(trimethylammonio)propyl]carbo diimide (I) has an acidic group of pK 6.8, and decomps. in its acid form as the dication by reaction with AcO- or H2O. Reaction of the carboxylate

anion with I is general-acid catalyzed, and the D2O solvent isotope

indicates a rate-detg. proton transfer except for the oxonium ion acting as acid. A mechanism involving proton transfer concerted with nucleophilic attack by AcO- is consistent with the weak basicity of the isourea adduct. The 3rd-order term involving HOAc, AcO- and carbodiimide carries .apprx.60% of the total reaction flux at pH 6.80 and 1 M total HOAc buffer concn. At this pH .apprx.40% of the reaction flux proceeds via a stepwise mechanism with specific acid catalysis. Intramol. general-acid catalysis occurs in the reaction of HO2CCEt2CO2- with I, and the effective molarity compared with intermol. catalysis is 15 M. Attack of carboxylate anions on I with N-(chloroethyl)morpholinium ion as the general acid has a Broensted-type .beta.N of 0.46.

IT 32897-26-0

RL: RCT (Reactant)

(reaction of, with phenethyl tosylate)

32897-26-0 CAPLUS

Urea, N-[3-(dimethylamino)propyl]-N'-ethyl- (9CI) (CA INDEX NAME) CN

EtNH-C-NH-(CH₂)₃-NMe₂

L12 ANSWER 15 OF 15 CAPLUS COPYRIGHT 2001 ACS

1981:135598 CAPLUS ACCESSION NUMBER:

94:135598 DOCUMENT NUMBER:

TITLE:

New immunochemical-glass conjugates

Sugiura, Masakazu; Kikutake, Junichiro; Yoshida, INVENTOR(S):

Masaru; Kondo, Shigeharu

Sanyo Chemical Industries, Ltd., Japan PATENT ASSIGNEE(S):

Fr. Demande, 30 pp. SOURCE:

CODEN: FRXXBL

Patent DOCUMENT TYPE:

French LANGUAGE: FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

APPLICATION NO. DATE KIND DATE PATENT NO. _____ ____ _____ FR 2435715 A1 19800404 FR 2435715 B1 19830708 FR 1979-2447 19790131 19800404

A method is described for the prepn. of a conjugate between a substance AΒ with immunol. activity (antigen or antibody) and frosted glass by using a silane coupling agent and, if necessary, a crosslinking agent. The frosted glass is reacted with a silane coupling agent which has an alkoxy silvl or halo silvl group which can react with a silanol group, as well

as a functional group (carboxyl, epoxy, aldehyde, etc.) which can react with amino, carboxyl, or thiol groups. The product is then reacted with the antigen or antibody in the presence of a crosslinking agent, when necessary. The crosslinking agent is an aliph. dialdehyde, a dichlorotriazine, a dimaleimide, or a maleimidocarboxyl-Nhydroxysuccinimide ester and can cause crosslinking between the amino,

```
carboxyl, or thiol groups of the silane and corresponding groups of the
    antigen or antibody. The antigen can be a hormone, protein, or an
    antigenic component of a pathogenic bacterium or virus or protozoan.
    Thus, ground-glass tubes were incubated with a soln. of 0.5%
     .gamma.-aminopropyl-triethoxysilane in Me2CO, followed by incubation at
    37.degree. for 2 h with a soln. contg. IgG and N-ethyl-N'-
    dimethylaminopropylcarbodiimide. Unconjugated proteins were washed out,
    and 63 .mu.g protein was fixed per g of glass. Glass beads can also be
    used, as for the detn. of insulin and .alpha.-fetoproteins by sandwich
    enzyme immunoassay.
ΙT
    32897-26-0
    RL: ANST (Analytical study)
        (in IgG immobilization on glass for immunoassay)
     32897-26-0 CAPLUS
RN
    Urea, N-[3-(dimethylamino)propyl]-N'-ethyl- (9CI) (CA INDEX NAME)
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EtNH-C-NH-(CH2)3-NMe2
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L2
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L3
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L4
          14060 S L4 FULL SUB=L3
L5
           5399 S L5 AND 3/N
L6
            734 S ETHYL(L) DIMETHYL(L) AMINO(L) PROPYL(L) UREA
L7
              0 S UREA, "N-ETHYL-N'-(3-DIMETHYLAMINOPROPYL)-"
rs
              4 S ETHYL(L) DIMETHYLAMINOPROPYL(L) UREA
L9
              3 S L9 AND 1/NC
L10
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L13
                 (L11 (L) THU/RL)
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L13 ANSWER 1 OF 2 CAPLUS COPYRIGHT 2001 ACS
                         2001:338479 CAPLUS
ACCESSION NUMBER:
                         Preparation of amides and ureas as activators of
TITLE:
                         soluble guanylate cyclase
```

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Selwood, David; Glen, Robert; Reynolds, Karen;
INVENTOR(S):
                        Wishart, Grant
                        University College London, UK
PATENT ASSIGNEE(S):
                         PCT Int. Appl., 101 pp.
SOURCE:
                        CODEN: PIXXD2
                        Patent
DOCUMENT TYPE:
                         English
LANGUAGE:
FAMILY ACC. NUM. COUNT:
                        1
PATENT INFORMATION:
                                          APPLICATION NO.
                                                           DATE
                     KIND DATE
     PATENT NO.
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                                                           20001106
                                          WO 2000-GB4249
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             BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                                                      A 19991105
                                        GB 1999-26286
PRIORITY APPLN. INFO.:
                                       US 2000-201382 P 20000502
     The title compds. R4PZNR1R2 [I; RA, R2 = alkyl; R1R2 together form
AB
     alkylene; Z = alkylene; P = a direct bond, X, Y, W, XY, YW, XYW (wherein
W
     = 0, S, NR3; R3 = H, alkyl; Y = iUV; V = a direct bond, alkylene; U = CS,
     CO, SO2, C(:NR); R = H, OH, alkyl; X = O, NR6; R6 = H, alkyl, alkenyl,
     etc.); R4 = alkyl, alkenyl, alkynyl, etc.], useful in the activation of
     sol. guanylate cyclase, were prepd. E.g., synthesis of the urea II,
     starting with 4-bromoaniline and 1-(3-aminopropyl)pyrrolidine, was given.
     Biol. data for compds. I (e.g., IC50 for inhibition of platelet
     aggregation) were presented.
     32897-26-0P
TΤ
     RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic
     preparation); THU (Therapeutic use); BIOL (Biological study);
     PREP (Preparation); USES (Uses)
        (prepn. of amides and ure as as activators of sol. guanylate cyclase)
RN
     32897-26-0 CAPLUS
     Urea, N-[3-(dimethylamino)propyl]-N'-ethyl- (9CI) (CA INDEX NAME)
CN
      0
EtNH-C-NH-(CH2)3-NMe2
REFERENCE COUNT:
                         (8) Catanese, B; BOLLETTINO CHIMICO FARMACEUTICO
REFERENCE(S):
1986,
                             V125(7), P228 CAPLUS
                         (9) Farbenfabriken Bayer Ag; DE 890958 C CAPLUS
                         (10) Glen, R; WO 0027394 A 2000 CAPLUS
                         (12) Hoechst Marion Roussel de Gmbh; EP 0908456 A
1999
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ALL CITATIONS AVAILABLE IN THE RE FORMAT
L13 ANSWER 2 OF 2 CAPLUS COPYRIGHT 2001 ACS
                               2000:725451 CAPLUS
ACCESSION NUMBER:
                               133:286497
DOCUMENT NUMBER:
                                Immunomodulatory compositions and methods of use
TITLE:
                                thereof
                                Onderdonk, Andrew B.; Tzianabos, Arthur O.; Miller,
INVENTOR(S):
                                Robert J.; Calias, Pericles
                                Genzyme Corporations, USA
PATENT ASSIGNEE(S):
                                PCT Int. Appl., 62 pp.
SOURCE:
                                CODEN: PIXXD2
                                Patent
DOCUMENT TYPE:
                                English
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
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                                   DATE
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      PATENT NO.
                           ____
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      WO 2000059490
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                SG, SI, SK, SL, TJ, TM, TR, TT, TZ, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
           RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF,
                CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                                                   us 1999-128177 P 19990406
PRIORITY APPLN. INFO.:
                                MARPATX 133:286497
OTHER SOURCE(S):
      The invention relates to immunomodulatory compns. and related methods.
      The immunomodulatory compns. are useful for the prevention of sepsis and
      the treatment and prevention of diseases absocd. with inflammation and/or
      NOS. CM-cellulose/N-ethyl-N'-(3-dimethylaminopropyl)urea formulations
are
      described.
IT
      32897-26<del>\</del>0
      RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
           (immundmodullatory compns.)
       32897-26-0\ CAPLUS
RN
      Urea, N-[3-\dimethylamino)propyl]-N'-ethyl- (9CI) (CA INDEX NAME)
CN
EtNH-C-NH-(CH2)3-NMè2
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CAPLUS

1999 CAPLUS

(13) Hoechst Marion Roussel de Gmbh; DE 19756388 A

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FULL ESTIMATED COST
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CA INDEXING COPYRIGHT (C) 2001 AMERICAN CHEMICAL SOCIETY (ACS)
FILE COVERS 1971 TO PATENT PUBLICATION DATE: 29 May 2001 (20010529/PD)
FILE LAST UPDATED: 29 May 2001 (20010529/ED)
HIGHEST PATENT NUMBER: US8411134
CA INDEXING IS CURRENT THROUGH 29 May 2001 (20010529/UPCA)
ISSUE CLASS FIELDS (/INCL) CURRENT THROUGH: 29 May 2001 (20010529/PD)
REVISED CLASS FIELDS (/NCL) LAST RELOADED: Apr 2001
USPTO MANUAL OF CLASSIFICATIONS THESAURUS ISSUE DATE: Apr 2001
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>>> Complete CA file indexing for chemical patents (or equivalents) <<<
>>> is included in file records. A thesaurus is available for the <<<
>>> USPTO Manual of Classifications in the /NCL, /INCL, and /RPCL
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>>> fields. This thesaurus includes catchword terms from the
                                                                     <<<
>>> USPTO/MOC subject headings and subheadings. Thesauri are also
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>>> available for the WIPO International Patent Classification
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>>> (IPC) Manuals, editions 1-6, in the /IC1, /IC2, /IC3, /IC4,
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>>> /IC5, and /IC (/IC6) fields, respectively. The thesauri in
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>>> the /IC5 and /IC fields include the corresponding catchword
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>>> terms from the IPC subject headings and subheadings.
This file contains CAS Registry Numbers for easy and accurate
substance identification.
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L2
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L3
                STRUCTURE UPLOADED
T.4
L5
          14060 S L4 FULL SUB=L3
L6
           5399 S L5 AND 3/N
            734 S ETHYL(L) DIMETHYL(L) AMINO(L) PROPYL(L) UREA
L7
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L8
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              3 S L9 AND 1/NC
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              1 S L10 AND 1/0
L11
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L12 15 S L11 L13 2 S L11/THU

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=> s 111 L14 2 L11

=> s 114 not 113

'THU' IS NOT A VALID CROSSOVER QUALIFIER FOR L11
Answer sets created in a different file may be field qualified with a limited set of qualifiers. Enter HELP CROSSOVER at an arrow prompt (=>) for specific information.

=> d ibib ab hitstr 1-2

L14 ANSWER 1 OF 2 USPATFULL

ACCESSION NUMBER: 96:29480 USPATFULL

TITLE: Non-specific reaction suppressor

INVENTOR(S): Ito, Michio, Indianapolis, IN, United States

Sugawa, Satoshi, Machida, Japan

Yanagida, Atsushi, Carmel, IN, United States

PATENT ASSIGNEE(S): Mitsubishi Kasei Corporation, Tokyo, Japan (non-U.S.

corporation)

NUMBER DATE

PATENT INFORMATION: US 5506151 19960409 APPLICATION INFO.: US 1994-194475 19940209 (8)

DOCUMENT TYPE: Utility

PRIMARY EXAMINER: Ceperley, Mary E.

LEGAL REPRESENTATIVE: Oblon, Spivak, McClelland, Maier & Neustadt

NUMBER OF CLAIMS: 16 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 13 Drawing Figure(s); 7 Drawing Page(s)

LINE COUNT: 575

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

nonspecific reaction suppressor)

AB A non-specific reaction suppressor for immunoassays having the formula: ##STR1## where R.sub.1, R.sub.2, Y, X, and R.sub.3 are defined in the specification.

IT 32897-26-0, 1-Ethyl3-(3-dimethylaminopropyl)urea (immunoassay uses latex particle-immobilized immunoreactant and

RN 32897-26-0 USPATFULL

CN Urea, N-[3-(dimethylamino)propyl]-N'-ethyl- (9CI) (CA INDEX NAME)

O || EtNH-C-NH-(CH₂)3-NMe₂

L14 ANSWER 2 OF 2 USPATFULL

ACCESSION NUMBER: 93:35827 USPATFULL

TITLE: Process for production of water-soluble carbodiimide

INVENTOR(S): Yoneyama, Takahiro, Matsudo, Japan

Odagiri, Masaki, Ushiku, Japan

Imanari, Makoto, Ami, Japan

PATENT ASSIGNEE(S): Research Association for Utilization of Light Oil,

Tokyo, Japan (non-U.S. corporation)

NUMBER DATE

PATENT INFORMATION: US 5208378 19930504

APPLICATION INFO.: US 1991-732123 19910718 (7)

NUMBER DATE

PRIORITY INFORMATION: JP 1990-189414 19900719

DOCUMENT TYPE: Utility

PRIMARY EXAMINER: Hollrah, Glennon H.
ASSISTANT EXAMINER: O'Sullivan, Peter G.
LEGAL REPRESENTATIVE: Wenderoth, Lind & Ponack

NUMBER OF CLAIMS: 10 EXEMPLARY CLAIM: 1 LINE COUNT: 239

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB A process for the production of a water-soluble carbodiimide, which comprises

- (1) allowing ethyl isothiocyanate to react with N,N-dimethyl-1,3propanediamine in an aromatic hydrocarbon solvent (first reaction step),
 - (2) removing hydrogen sulfide from a thiourea derivative formed in the first reaction step upon adding a hydrogen sulfide removing agent without isolating the thiourea derivative (second reaction step), and
 - (3) recovering a water-soluble carbodiimide from the resulting reaction mixture.

IT 32897-26-0P

(prepn. and dehydrosulfurization of)

RN 32897-26-0 USPATFULL

CN Urea, N-[3-(dimethylamino)propyl]-N'-ethyl- (9CI) (CA INDEX NAME)

O || EtNH-C-NH-(CH₂)₃-NMe₂

=> d ibib ab hitstr 1-2

L14 ANSWER 1 OF 2 USPATFULL

ACCESSION NUMBER: 96:29480 USPATFULL

TITLE: Non-specific reaction suppressor

INVENTOR(S): Ito, Michio, Indianapolis, IN, United States

Sugawa, Satoshi, Machida, Japan

Yanagida, Atsushi, Carmel, IN, United States

PATENT ASSIGNEE(S): Mitsubishi Kasei Corporation, Tokyo, Japan (non-U.S.

DATE

corporation)

-----PATENT INFORMATION: US 5506151 US 1994-194475 19940209 (8)

DOCUMENT TYPE:

PRIMARY EXAMINER: Ceperley, Mary E.

LEGAL REPRESENTATIVE: Oblon, Spivak, McClelland, Maier & Neustadt

NUMBER

NUMBER OF CLAIMS: 16 1 EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 13 Drawing Figure(s); 7 Drawing Page(s)

575 LINE COUNT:

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

A non-specific reaction suppressor for immunoassays having the formula: ##STR1## where R.sub.1, R.sub.2, Y, X, and R.sub.3 are defined in the specification.

IT 32897-26-0, 1-Ethyl3-(3-dimethylaminopropyl)urea

(immunoassay uses latex particle-immobilized immunoreactant and nonspecific reaction suppressor)

RN 32897-26-0 USPATFULL

Urea, N-[3-(dimethylamino)propyl]-N'-ethyl- (9CI) (CA INDEX NAME) CN

0 EtNH-C-NH-(CH₂)₃-NMe₂

L14 ANSWER 2 OF 2 USPATFULL

ACCESSION NUMBER: 93:35827 USPATFULL

Process for production of water-soluble carbodiimide TITLE: INVENTOR(S):

Yoneyama, Takahiro, Matsudo, Japan Odagiri, Masaki, Ushiku, Japan Imanari, Makoto, Ami, Japan

Research Association for Utilization of Light Oil, PATENT ASSIGNEE(S):

Tokyo, Japan (non-U.S. corporation)

NUMBER ______ US 5208378 19930504 PATENT INFORMATION: US 1991-732123 19910718 (7) APPLICATION INFO.:

NUMBER DATE

PRIORITY INFORMATION: JP 1990-189414 19900719

DOCUMENT TYPE: Utility

PRIMARY EXAMINER: Hollrah, Glennon H. ASSISTANT EXAMINER: O'Sullivan, Peter G. Wenderoth, Lind & Ponack LEGAL REPRESENTATIVE:

NUMBER OF CLAIMS: 10 EXEMPLARY CLAIM: 1 LINE COUNT: 239

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

- AB A process for the production of a water-soluble carbodiimide, which comprises
- (1) allowing ethyl isothiocyanate to react with N,N-dimethyl-1,3propanediamine in an aromatic hydrocarbon solvent (first reaction step),
 - (2) removing hydrogen sulfide from a thiourea derivative formed in the first reaction step upon adding a hydrogen sulfide removing agent without isolating the thiourea derivative (second reaction step), and
 - (3) recovering a water-soluble carbodiimide from the resulting reaction mixture.

IT 32897-26-0P

(prepn. and dehydrosulfurization of)

RN 32897-26-0 USPATFULL

CN Urea, N-[3-(dimethylamino)propyl]-N'-ethyl- (9CI) (CA INDEX NAME)

=> file marpat COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION FULL ESTIMATED COST 22.20 363.62 DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) SINCE FILE TOTAL ENTRY SESSION CA SUBSCRIBER PRICE 0.00 -10.00

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FILE CONTENT: 1988-PRESENT (VOL 104 ISS 15-VOL 134 ISS 22) (20010525/ED)

MOST RECENT CITATIONS FOR PATENTS FROM FIVE MAJOR ISSUING AGENCIES (COVERAGE TO THESE DATES IS NOT COMPLETE):

US 6225295 01 MAY 2001 DE 10035614 26 APR 2001 EP 1095952 02 MAY 2001 JP 200112290 08 MAY 2001 WO 200103079 03 MAY 2001

MARPAT structure search limits have been raised. Enter HELP SLIMIT for details.

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     FILE 'REGISTRY' ENTERED AT 10:38:44 ON 01 JUN 2001
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L3
          36270 S L1 FULL
L4
                STRUCTURE UPLOADED
L5
          14060 S L4 FULL SUB=L3
L6
           5399 S L5 AND 3/N
L7
            734 S ETHYL(L) DIMETHYL(L) AMINO(L) PROPYL(L) UREA
              0 S UREA, "N-ETHYL-N'-(3-DIMETHYLAMINOPROPYL)-"
rs
              4 S ETHYL(L) DIMETHYLAMINOPROPYL(L) UREA
L9
L10
              3 S L9 AND 1/NC
              1 S L10 AND 1/0
L11
     FILE 'CAPLUS' ENTERED AT 10:52:57 ON 01 JUN 2001
L12
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L13
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     FILE 'USPATFULL' ENTERED AT 10:55:19 ON 01 JUN 2001
L14
              2 S L11
     FILE 'MARPAT' ENTERED AT 10:56:49 ON 01 JUN 2001
=> s 111
'NC' IS NOT A VALID FIELD CODE
'O' IS NOT A VALID FIELD CODE
             7 ETHYL
             0 DIMETHYLAMINOPROPYL
             0 UREA
             1 UREAS
             1 UREA
                 (UREA OR UREAS)
             0 ETHYL(L)DIMETHYLAMINOPROPYL(L)UREA
             0 1/NC
             0 1/0
L15
             0 L10 AND 1/O
=> s 19
             7 ETHYL
             0 DIMETHYLAMINOPROPYL
             0 UREA
             1 UREAS
             1 UREA
                 (UREA OR UREAS)
L16
             0 ETHYL(L) DIMETHYLAMINOPROPYL(L) UREA
=> file beil
COST IN U.S. DOLLARS
                                                  SINCE FILE
                                                                   TOTAL
                                                       ENTRY
                                                                 SESSION
FULL ESTIMATED COST
                                                        9.19
                                                                  372.81
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)
                                                  SINCE FILE
                                                                   TOTAL
                                                       ENTRY
                                                                 SESSION
CA SUBSCRIBER PRICE
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0.00

-10.00

FILE 'BEILSTEIN' ENTERED AT 10:57:15 ON 01 JUN 2001 COPYRIGHT (c) 2001 Beilstein-Institut zur Foerderung der Chemischen Wissenschaften licensed to Beilstein Chemiedaten & Software GmbH and MDL Information Systems GmbH FILE LAST UPDATED: 6 MAR 2000 FILE COVERS 1779 TO 2000. *** CÁS REGISTRY NUMBERS FOR 4,356,237 SUBSTANCES AVAILABLE *** *** FILE CONTAINS 7,688,486 SUBSTANCES *** ******************* * PLEASE NOTE THAT THERE ARE NO FORMATS FREE OF COST. * SET NOTICE FEATURE: THE COST ESTIMATES CALCULATED FOR SET NOTICE * ARE BASED ON THE HIGHEST PRICE CATEGORY. THEREFORE; THESE * ESTIMATES MAY NOT REFLECT THE ACTUAL COSTS. * FOR PRICE INFORMATION SEE HELP COST ***************** => d his (FILE 'HOME' ENTERED AT 10:38:39 ON 01 JUN 2001) FILE 'REGISTRY' ENTERED AT 10:38:44 ON 01 JUN 2001 STRUCTURE UPLOADED L1L2 50 S L1 L3 36270 S L1 FULL STRUCTURE UPLOADED L4L5 14060 S L4 FULL SUB=L3 L6 5399 S L5 AND 3/N 734 S ETHYL(L) DIMETHYL(L) AMINO(L) PROPYL(L) UREA L7 0 S UREA, "N-ETHYL-N'-(3-DIMETHYLAMINOPROPYL)-" L8 L9 4 S ETHYL(L) DIMETHYLAMINOPROPYL(L) UREA L10 3 S L9 AND 1/NC L11 1 S L10 AND 1/0 FILE 'CAPLUS' ENTERED AT 10:52:57 ON 01 JUN 2001 L1215 S L11 L13 2 S L11/THU FILE 'USPATFULL' ENTERED AT 10:55:19 ON 01 JUN 2001 L14 2 S L11 FILE 'MARPAT' ENTERED AT 10:56:49 ON 01 JUN 2001 0 S L11 L15 0 S L9 L16

FILE 'BEILSTEIN' ENTERED AT 10:57:15 ON 01 JUN 2001

=> s 111 L17 1 L11

=> d all

L17 ANSWER 1 OF 1 COPYRIGHT 2001 BEILSTEIN CDS MDLI

Beilstein Reg. No. (BRN): 4961600 Beilstein

Molecular Formula (MF): C8 H19 N3 O

Autonom Name (AUN): 1-(3-dimethylamino-propyl)-3-ethyl-urea

Beilstein Reference (SO): 6-04

CAS Reg. No. (RN): 32897-26-0
Beilstein Pref. RN (BPR): 32897-26-0
Formula Weight (FW): 173.26

Lawson Number (LN): 3027; 2826; 2817; 1762

Ring System Data:

Number of Rings (CNR): 0
Acyclic Heteros (CNAH): 4

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Preparation:

PRE

Start: BRN=773743 isocyanatoethane, BRN=605293 N,N-dimethyl-propane-1,3-

diamine

Time: 2 hour(s)
Solv: diethyl ether
Ambient Temperature

Reference(s):

1. Williams, Andrew; Ibrahim, Ibrahim T., J.Amer.Chem.Soc., 103 <1981>

24,

7090-7095, LA: EN, CODEN: JACSAT

Note(s):

2. Yield given

=> file reg

COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION FULL ESTIMATED COST 14.34 387.15 DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) SINCE FILE TOTAL ENTRY SESSION CA SUBSCRIBER PRICE 0.00 -10.00

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STRUCTURE FILE UPDATES: 30 MAY 2001 HIGHEST RN 339046-06-9 DICTIONARY FILE UPDATES: 30 MAY 2001 HIGHEST RN 339046-06-9

TSCA INFORMATION NOW CURRENT THROUGH January 11, 2001

Please note that search-term pricing does apply when conducting SmartSELECT searches.

Structure search limits have been increased. See HELP SLIMIT for details.

=> d his

(FILE 'HOME' ENTERED AT 10:38:39 ON 01 JUN 2001)

FILE 'REGISTRY' ENTERED AT 10:38:44 ON 01 JUN 2001 L1STRUCTURE UPLOADED L2 50 S L1 36270 S L1 FULL L3 L4 STRUCTURE UPLOADED L5 14060 S L4 FULL SUB=L3 L6 5399 S L5 AND 3/N 734 S ETHYL (L) DIMETHYL (L) AMINO (L) PROPYL (L) UREA L7 0 S UREA, "N-ETHYL-N'-(3-DIMETHYLAMINOPROPYL)-" 4 S ETHYL(L)DIMETHYLAMINOPROPYL(L)UREA L9 L10 3 S L9 AND 1/NC L11 1 S L10 AND 1/0

FILE 'CAPLUS' ENTERED AT 10:52:57 ON 01 JUN 2001 15 S L11

L12 15 S L11 L13 2 S L11/THU

FILE 'USPATFULL' ENTERED AT 10:55:19 ON 01 JUN 2001

L14 2 S L11

FILE 'MARPAT' ENTERED AT 10:56:49 ON 01 JUN 2001

L15 0 S L11 L16 0 S L9

FILE 'REGISTRY' ENTERED AT 10:57:49 ON 01 JUN 2001

=> Uploading 489.str

L18 STRUCTURE UPLOADED

=> s 118 sub=13 full FULL SUBSET SEARCH INITIATED 11:00:01 FILE 'REGISTRY' FULL SUBSET SCREEN SEARCH COMPLETED - 32305 TO ITERATE 100.0% PROCESSED 32305 ITERATIONS SEARCH TIME: 00.00.05

11247 ANSWERS

L19 11247 SEA SUB=L3 SSS FUL L18

=> d scan

L19 11247 ANSWERS REGISTRY COPYRIGHT 2001 ACS

IN Urea, N-[2-[[[(4-ethoxyphenyl)amino]carbonyl]amino]ethyl]-N-(4phenoxyphenyl)-N'-phenyl- (9CI)

MF C30 H30 N4 O4

$$\begin{array}{c|c} \text{PhO} & \begin{array}{c|c} \text{O} \\ \parallel \\ \text{C-NHPh} \\ \text{N-CH}_2\text{-CH}_2\text{-NH-C-NH} \end{array} \end{array} \text{OEt}$$

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):1

L19 11247 ANSWERS REGISTRY COPYRIGHT 2001 ACS

IN Urea, N'-(2-fluorophenyl)-N-(4-fluorophenyl)-N-[2-[[[(2-methoxyphenyl)amino]carbonyl]amino]ethyl]- (9CI)

MF C23 H22 F2 N4 O3

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

=> d his

(FILE 'HOME' ENTERED AT 10:38:39 ON 01 JUN 2001)

FILE 'REGISTRY' ENTERED AT 10:38:44 ON 01 JUN 2001
L1 STRUCTURE UPLOADED
L2 50 S L1
L3 36270 S L1 FULL
L4 STRUCTURE UPLOADED
L5 14060 S L4 FULL SUB=L3
L6 5399 S L5 AND 3/N
L7 734 S ETHYL(L) DIMETHYL(L) AMINO(L) PROPYL(L) UREA

```
0 S UREA, "N-ETHYL-N'-(3-DIMETHYLAMINOPROPYL)-"
^{18}
              4 S ETHYL(L) DIMETHYLAMINOPROPYL(L) UREA
L9
              3 S L9 AND 1/NC
L10
L11
              1 S L10 AND 1/0
     FILE 'CAPLUS' ENTERED AT 10:52:57 ON 01 JUN 2001
L12
             15 S L11
L13
              2 S L11/THU
     FILE 'USPATFULL' ENTERED AT 10:55:19 ON 01 JUN 2001
L14
              2 S L11
     FILE 'MARPAT' ENTERED AT 10:56:49 ON 01 JUN 2001
L15
              0 S L11
              0 S L9
L16
     FILE 'BEILSTEIN' ENTERED AT 10:57:15 ON 01 JUN 2001
L17
              1 S L11
     FILE 'REGISTRY' ENTERED AT 10:57:49 ON 01 JUN 2001
                STRUCTURE UPLOADED
L18
          11247 S L18 FULL SUB=L3
L19
=> s 118
SAMPLE SEARCH INITIATED 11:00:39 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 2732 TO ITERATE
 36.6% PROCESSED
                    1000 ITERATIONS
                                                               50 ANSWERS
INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)
SEARCH TIME: 00.00.02
FULL FILE PROJECTIONS:
                        ONLINE **COMPLETE**
                                **COMPLETE**
                        BATCH
PROJECTED ITERATIONS:
                             51507 TO
                                         57773
PROJECTED ANSWERS:
                             10242 TO
                                          13142
             50 SEA SSS SAM L18
L20
=> d scan
L20 50 ANSWERS REGISTRY COPYRIGHT 2001 ACS
     Benzamide, 2.5-bis (2.2.2-trifluoroethoxy)-N-[2-[[[3-
     (trifluoromethyl)phenyl]amino]carbonyl]amino]ethyl]- (9CI)
MF
     C21 H18 F9 N3 O4
```

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

=> s 118 css

SAMPLE SEARCH INITIATED 11:01:00 FILE 'REGISTRY' SAMPLE SCREEN SEARCH COMPLETED - 2732 TO ITERATE

36.6% PROCESSED 1000 ITERATIONS INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED) SEARCH TIME: 00.00.01

ATIONS 0 ANSWERS

0

FULL FILE PROJECTIONS: ONLINE **COMPLETE**

BATCH **COMPLETE**

PROJECTED ITERATIONS: 51507 TO 57773

PROJECTED ANSWERS: 0 TO

L21 0 SEA CSS SAM L18

=> s 118 css full FULL SEARCH INITIATED 11:01:07 FILE 'REGISTRY' FULL SCREEN SEARCH COMPLETED - 54245 TO ITERATE

100.0% PROCESSED 54245 ITERATIONS SEARCH TIME: 00.00.04

8 ANSWERS

L22 8 SEA CSS FUL L18

=> d scan

L22 8 ANSWERS REGISTRY COPYRIGHT 2001 ACS
IN Urea, N-[4-(dimethylamino)butyl]-N'-methyl- (9CI)
MF C8 H19 N3 O

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):5

L22 8 ANSWERS REGISTRY COPYRIGHT 2001 ACS

IN Urea, N-[2-(dimethylamino)ethyl]-N'-methyl- (9CI)

MF C6 H15 N3 O

CI COM

L22 8 ANSWERS REGISTRY COPYRIGHT 2001 ACS IN Urea, N-(3-aminopropyl)-N'-methyl- (9CI) MF C5 H13 N3 O

L22 8 ANSWERS REGISTRY COPYRIGHT 2001 ACS
IN Urea, N-[3-(dimethylamino)propyl]-N'-methyl- (9CI)
MF C7 H17 N3 O

$$\begin{array}{c} \text{O} \\ \parallel \\ \text{MeNH-C-NH- (CH2)}_{3} - \text{NMe}_{2} \end{array}$$

L22 8 ANSWERS REGISTRY COPYRIGHT 2001 ACS
IN Urea, N-(2-aminoethyl)-N'-methyl-, compd. with 2,4,6-trinitrophenol (1:1) (9CI)
MF C6 H3 N3 O7 . C4 H11 N3 O

CM 1

$$\begin{tabular}{l} O \\ || \\ MeNH-C-NH-CH_2-CH_2-NH_2 \\ \end{tabular}$$

CM 2

L22 8 ANSWERS REGISTRY COPYRIGHT 2001 ACS
IN Urea, N-(2-aminoethyl)-N'-methyl-, monohydrochloride (9CI)
MF C4 H11 N3 O . Cl H

 $\begin{tabular}{ll} \tt O & & & & \\ \parallel & & & \\ \tt MeNH-C-NH-CH_2-CH_2-NH_2 \end{tabular}$

● HCl

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

=> file caplus SINCE FILE COST IN U.S. DOLLARS TOTAL ENTRY SESSION FULL ESTIMATED COST 166.92 554.07 SINCE FILE TOTAL DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) ENTRY SESSION 0.00 -10.00CA SUBSCRIBER PRICE

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FILE COVERS 1947 - 1 Jun 2001 VOL 134 ISS 23 FILE LAST UPDATED: 30 May 2001 (20010530/ED)

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INVENTOR(S):

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(FILE 'HOME' ENTERED AT 10:38:39 ON 01 JUN 2001)
     FILE 'REGISTRY' ENTERED AT 10:38:44 ON 01 JUN 2001
                STRUCTURE UPLOADED
L1
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          36270 S L1 FULL
L4
                STRUCTURE UPLOADED
          14060 S L4 FULL SUB=L3
L5
L6
           5399 S L5 AND 3/N
L7
            734 S ETHYL(L) DIMETHYL(L) AMINO(L) PROPYL(L) UREA
              0 S UREA, "N-ETHYL-N'-(3-DIMETHYLAMINOPROPYL)-"
rac{1}{8}
              4 S ETHYL(L) DIMETHYLAMINOPROPYL(L) UREA
L9
L10
              3 S L9 AND 1/NC
              1 S L10 AND 1/0
L11
     FILE 'CAPLUS' ENTERED AT 10:52:57 ON 01 JUN 2001
L12
             15 S L11
              2 S L11/THU
L13
     FILE 'USPATFULL' ENTERED AT 10:55:19 ON 01 JUN 2001
              2 S L11
L14
     FILE 'MARPAT' ENTERED AT 10:56:49 ON 01 JUN 2001
L15
              0 S L11
L16
              0 S L9
     FILE 'BEILSTEIN' ENTERED AT 10:57:15 ON 01 JUN 2001
L17
              1 S L11
     FILE 'REGISTRY' ENTERED AT 10:57:49 ON 01 JUN 2001
                STRUCTURE UPLOADED
L18
          11247 S L18 FULL SUB=L3
L19
             50 S L18
L20
L21
              0 S L18 CSS
L22
              8 S L18 CSS FULL
     FILE 'CAPLUS' ENTERED AT 11:01:43 ON 01 JUN 2001
=> s 122
            10 L22
L23
=> d ibib ab hitstr 1-10
L23 ANSWER 1 OF 10 CAPLUS COPYRIGHT 2001 ACS
                         1990:118630 CAPLUS
ACCESSION NUMBER:
                         112:118630
DOCUMENT NUMBER:
                         3-Amino-2-hydroxypropyl furoates or
TITLE:
                         thiophenecarboxylates as .beta.-adrenergic blockers
```

Kam, Sheung T.; Matier, William L.; Patil, Ghanshyam;

Mai, Khuong H. X.

du Pont de Nemours, E. I., and Co., USA PATENT ASSIGNEE(S):

SOURCE:

U.S., 20 pp. Cont.-in-part of U.S. 4,582,855.

CODEN: USXXAM

DOCUMENT TYPE:

Patent English

LANGUAGE: 2

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PAT	ENT NO.		KIND	DATE		APP	LICATION NO.	DATE
US	4798892		Α	19890117		US	1986-851629	19860414
US	4582855		Α	19860415		US	1981-320773	19811112
ZA	8207749		A	19830831		ZA	1982-7749	19821022
EP	93765		A1	19831116		EP	1982-903569	19821028
EP	93765		В1	19861210				
	R: BE,	CH, D	E, FR,	, GB, LI,	LU, N	NL, S	E	
CA	1201438		A1	19860304		CA	1982-415282	19821110
ES	517296		A1	19831201		ES	1982-517296	19821111
${\tt IL}$	67243		A1	19870331		IL	1982-67243	19821112
ES	523804		A1	19841101		ES	1983-523804	19830701
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NO	8302526		A	19830711		NO	1983-2526	19830711
NO	170926		В	19920921				
NO	170926		С	19921230				
ES	530788		A1	19850601		ES	1984-530788	19840320
US	4810717		A	19890307		US	1986-838082	19860310
US	4935421		Α	19900619		US	1989-318147	19890301
PRIORITY	APPLN.	<pre>INFO.:</pre>			US	5 198	1-320773	19811112
					US	5 198	6-838082	19860310

MARPAT 112:118630 OTHER SOURCE(S):

Title compds. ArCO2CH2CH(OH)CH2NHWNR1B [I; Ar = (substituted) furyl or thienyl or Ph; W = C1-10 alkylene; B = COR2, CONR2R3, SO2R2, SO2NR2R3, CO2R2; R1, R2, R3 = H, alkyl, alkoxyalkyl, alkenyl, Ph, etc.; R2R3N = morpholino; R2 .noteq. H when B = SO2R2, CO2R2] are prepd. To a soln. of glycidol and Et3N in Et2O was added 2-furoyl chloride, giving 88.0% of ester II, which in DMF was heated with 1,1-dimethyl-2-[(morpholinocarbonyl)amino]ethylamine at 70.degree. to give furoate ester III, isolated as its oxalate (23.0%). III at 2.4 .mu.g/kg/min gave 40% inhibition of heart rate response to isoproterenol in anesthetized dogs, and exhibited pA2 (guinea pig atria in vitro) of 7.6.

122036-80-0P IT

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn. and reaction of, in prepn. of .beta.-adrenergic blockers)

122036-80-0 CAPLUS

Urea, N-(2-aminoethyl)-N'-methyl-, monohydrochloride (9CI) (CA INDEX NAME)

MeNH-C-NH-CH2-CH2-NH2

O || MeNH-C-NH-CH₂-CH₂-NH₂

⊕ HCl

L23 ANSWER 2 OF 10 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 1989:211700 CAPLUS

DOCUMENT NUMBER: 110:211700

TITLE: Preparation of carbodiimides by a phase-transfer

catalytic method

AUTHOR(S): Jaszay, Zsuzsa M.; Petnehazy, Imre; Toke, Laszlo;

Szajani, Bela

CORPORATE SOURCE: Szerv. Kem. Technol. Tansz., Budapesti Muszaki

Egyet.,

Budapest, 1521, Hung.

SOURCE: Magy. Kem. Foly. (1988), 94(6-7), 246-9

CODEN: MGKFA3; ISSN: 0025-0155

DOCUMENT TYPE: Journal LANGUAGE: Hungarian

AB A new method is described for the prepn. of carbodismides by dehydration

of ureas with arom. sulfonic acid chloride under solid-liq.

phase-transfer

catalytic conditions using solid K2CO3 as base and a lipophile quaternary ammonium salt as catalyst. The method is generally applicable for the synthesis of disubstituted carbodiimides, but esp. useful for unsym. substituted carbodiimides. Most of the carbodiimides prepd. have been identified in the form of the more stable, cryst. quaternary salt.

IT 111681-36-8

RL: RCT (Reactant)

(dehydration of, with arenesulfonyl chloride under phase-transfer catalytic conditions)

RN 111681-36-8 CAPLUS

CN Urea, N-[3-(dimethylamino)propyl]-N'-methyl- (9CI) (CA INDEX NAME)

O || MeNH-C-NH-(CH₂)₃-NMe₂

L23 ANSWER 3 OF 10 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 1988:68296 CAPLUS

DOCUMENT NUMBER: 108:68296

TITLE: Choline-like nitrosoalkylurea derivatives and their

antitumor activity

AUTHOR(S): Belyaev, A. A.; Radina, L. B.; Anoshina, G. M.;

Peretolchina, N. M.; Sof'ina, Z. P.

CORPORATE SOURCE: Inst. Khim., Sverdlovsk, USSR

SOURCE: Khim.-Farm. Zh. (1987), 21(8), 940-5

CODEN: KHFZAN; ISSN: 0023-1134

DOCUMENT TYPE:

Journal

LANGUAGE:

Russian

N,N-Dimethylpropanediamine, N,N-dimethylbutanediamine, and N,N,N'-trimethylethylenediamine were carbamoylated with suitable alkyl isocyanates, the urea derivs. formed were quaternized with Me tosylate, and the quaternized derivs. were treated with N2O3 to give nitrosoalkyl urea derivs., R1R2NCONR(CH2)nN+Me3 TsO- (R = H, Me or NO, R1 = Me, CH2CH2Cl2, or cyclohexyl and R2 = H or NO, and n = 2-4). The antitumor activity and toxicity of these compds. were evaluated. Toxicity of the disubstituted nitrosoalkylureas in comparison with choline-like nitrosoalkylureas was maintained at max. tolerable dose, 10-30 mg/kg, while that of the trisubstituted derivs. it decreased to the max. tolerable dose of 250-300 mg/kg. C1CH2CH2N(NO)CONMe(CH2)2N+Me TsO-showed

the highest antitumor activity at 250 mg/kg. Structure-activity relations

are discussed.

IT 111681-36-8P 112557-32-1P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn. and quaternization of)

RN 111681-36-8 CAPLUS

CN Urea, N-[3-(dimethylamino)propyl]-N'-methyl- (9CI) (CA INDEX NAME)

RN 112557-32-1 CAPLUS

CN Urea, N-[4-(dimethylamino)butyl]-N'-methyl- (9CI) (CA INDEX NAME)

O || MeNH-C-NH-(CH₂)₄-NMe₂

L23 ANSWER 4 OF 10 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 1988:5307 CAPLUS

DOCUMENT NUMBER: 108:5307

TITLE: Preparation of carbodiimides using phase-transfer

catalysis

AUTHOR(S): Jaszay, Zsuzsa M.; Petnehazy, Imre; Toke, Laszlo;

Szajani, Bela

CORPORATE SOURCE: Tech. Univ. Budapest, Badapest, H-1521, Hung.

SOURCE: Synthesis (1987), (5), 520-3

CODEN: SYNTBF; ISSN: 0039-7881

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 108:5307

AB RN:C:NR1 (R = cyclohexyl, Ph, Bu, Me, Me3C; R1 = aminoalkyl, PhCH2, cyclohexyl, Me3C) were prepd. by dehydration of ureas with arenesulfonyl chlorides under solid-liq. phase-transfer conditions with solid K2CO3 as base and PhCH2N+Et3 Cl- as catalyst. The method was esp. useful for the

INDEX NAME)

```
synthesis of unsym. substituted carbodiimides. The basic carbodiimides
     were converted into more stable, cryst. quaternary salts.
ΙT
     111681-36-8
     RL: RCT (Reactant)
        (dehydration of, by arylsulfonyl chloride)
     111681-36-8 CAPLUS
RN
     Urea, N-[3-(dimethylamino)propyl]-N'-methyl- (9CI) (CA INDEX NAME)
CN
     0
MeNH-C-NH-(CH<sub>2</sub>)<sub>3</sub>-NMe<sub>2</sub>
L23 ANSWER 5 OF 10 CAPLUS COPYRIGHT 2001 ACS
                         1987:32335 CAPLUS
ACCESSION NUMBER:
DOCUMENT NUMBER:
                         106:32335
                         Nitrosoalkylureas based on alkylammonium salts and
TITLE:
                         their antitumor activity
                         Belyaev, A. A.; Gopko, V. F.; Radina, L. B.;
AUTHOR(S):
                         Peretolchina, N. M.; Sof'ina, Z. P.; Anoshina, G. M.;
                         Zubova, T. E.
                         Inst. Khim., Sverdlovsk, USSR
CORPORATE SOURCE:
                         Khim.-Farm. Zh. (1986), 20(5), 532-6
SOURCE:
                         CODEN: KHFZAN; ISSN: 0023-1134
DOCUMENT TYPE:
                         Journal
LANGUAGE:
                         Russian
     Seven title compds. were prepd. by reaction of dimethyl(aminoethyl)amine
     with the appropriate isocyanate, followed by either quaternization or
     hydrochloride formation. In vitro tests of neoplasm inhibition showed 2
     chloroethyl derivs. to be the most potent. Given i.p. to mice bearing
     various tumors, the hydrochloride form was more active and more toxic
than
     the quaternary salt form. Structure activity relations are discussed.
     105996-25-6P 105996-27-8P
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (prepn. of)
RN
     105996-25-6 CAPLUS
     Urea, N-[2-(dimethylamino)ethyl]-N'-methyl- (9CI) (CA INDEX NAME)
CN
      0
MeNH-C-NH-CH2-CH2-NMe2
RN
     105996-27-8 CAPLUS
     Urea, N-[2-(dimethylamino)ethyl]-N'-methyl-, monohydrochloride (9CI) (CA
CN
```

⊕ HCl

L23 ANSWER 6 OF 10 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 1984:6088 CAPLUS

DOCUMENT NUMBER: 100:6088

TITLE: 2-Hydroxypropylamine aryl ester derivatives

INVENTOR(S): Kam, Sheung Tsam; Matier, William L. PATENT ASSIGNEE(S): American Hospital Supply Corp., USA

SOURCE: PCT Int. Appl., 70 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PAT	TENT NO.		KIND	DATE		APPLICATION NO.	DATE
WO				19830526 , RO, SU		WO 1982-US1536	19821028
				, FR, GB,	LU,	NL, SE	
US						US 1981-320773	19811112
						ZA 1982-7749	
	8210120		A1	19830601		AU 1982-10120	19821028
	93765		A1	19831116		EP 1982-903569	19821028
ΕP	93765		B1	19861210			
	R: BE,	CH,	DE, FR	, GB, LI,	LU,	NL, SE	
	562862		B2			AU 1983-10120	19821028
CA	1201438		A1	19860304		CA 1982-415282	19821110
ES	517296		A1	19831201		ES 1982-517296	19821111
ΙL	67243		A1	19870331		IL 1982-67243	19821112
ES	523804		A 1			ES 1983-523804	
ES	523805		A 1	19841101		ES 1983-523805	19830701
NO	8302526					NO 1983-2526	19830711
NO	170926		В	19920921			
NO	170926			19921230			
	58501724		T 2	19831013		JP 1982-503552	19830712
JP	63020424		B4	19880427			
ES	530788		A 1	19850601		ES 1984-530788	19840320
US	4810717		Α	19890307		US 1986-838082	19860310
US	4935421		A	19900619		US 1989-318147	
RITY	APPLN.	INFO	. :			us 1981-320773	
						WO 1982-US1536	
						US 1986-838082	

AB .beta.-Blockers RCO2CH2CH(OH)CH2NH-X-R1 [R = (un)substituted aryl, heterocyclic; X = C1-C10 alkylene; R1 = NR2COR3, NR2CONR3R4, NR2SO2R3, NR2SO2NR3R4, NR2CO2R3; R2-R4 = H, alkyl, alkoxyalkyl, cycloalkyl, alkenyl,

alkynyl, aryl, heteroaryl, aralkyl; NR3R4 = 5-7 membered heterocycle]
were

prepd. Thus EtOAc reacted with H2NCH2CMe2NH2 to give 57.4% AcNHCH2CMe2NH2

(I). 2-FC6H4COC1 reacted with glycidol to give 2-FC6H4CO2R5 (R5 = 2,3-epoxypropyl), which was treated with I to give AcNHCH2CMe2NHCH2CH(OH)CH2O2CC6H4F-4 (II). At 2.7 mg/kg II 3 h after administration gave 61% inhibition of heart rate response to

isoproterenol

in dogs. The aryl esters of this invention were also useful in the treatment of glaucoma (no data).

ΙT 75930-29-9P

RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. of)

75930-29-9 CAPLUS RN

Urea, N-(2-aminoethyl)-N'-methyl- (9CI) (CA INDEX NAME) CN

0 MeNH-C-NH-CH2-CH2-NH2

L23 ANSWER 7 OF 10 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

1984:6069 CAPLUS

DOCUMENT NUMBER:

100:6069

TITLE:

P-Substituted 3-phenoxy-1-ureidoalkylamino-2-propanols

INVENTOR(S):

Gustafsson, Bill Bemjamin Rudolf; Hedberg, Sven

Anders; Lundgren, Bo Torsten

PATENT ASSIGNEE(S):

SOURCE:

Hassle AB, Swed.

Brit. UK Pat. Appl., 25 pp.

CODEN: BAXXDU

DOCUMENT TYPE:

LANGUAGE:

Patent English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
GB 2111500	A1	19830706	GB 1982-35707	19821215
GB 2111500	В2	19850807		
EP 85286	A1	19830810	EP 1982-850257	19821210
EP 85286	B1	19860430		
R: AT, BE,	CH, DE	, FR, GB, I	T, LI, LU, NL, SE	
AT 19510	E	19860515	AT 1982-850257	19821210
ZA 8209249	Α	19830928	ZA 1982-9249	19821215
FI 8204339	Α	19830618	FI 1982-4339	19821216
NO 8204237	Α	19830620	NO 1982-4237	19821216
NO 154835	В	19860922		
NO 154835	С	19870102		
JP 58110556	A2	19830701	JP 1982-219367	19821216
ES 518268	A1	19840216	ES 1982-518268	19821216
HU 31106	0	19840428	HU 1982-4066	19821216
CA 1178588	A1	19841127	CA 1982-417848	19821216
RO 87523	В3	19850831	RO 1982-109344	19821216
CS 239948	В2	19860116	CS 1982-8725	19821216

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19860116
                                            CS 1982-9249
                                                              19821216
     CS 239935
                       B2
                             19830618
                                            DK 1982-5601
                                                              19821217
     DK 8205601
                       Α
     AU 8291637
                       A1
                             19830623
                                            AU 1982-91637
                                                              19821217
                       A5
                             19840411
                                            DD 1982-246176
                                                              19821217
     DD 208799
                       В1
                             19870228
                                            PL 1982-239591
                                                              19821217
     PL 139609
     SU 1169533
                       A3
                             19850723
                                            SU 1983-3657075
                                                              19831031
                                            ES 1983-527314
                                                              19831116
     ES 527314
                       A1
                             19841201
                                                              19840901
                       A1
                             19851201
                                            ES 1984-535597
     ES 535597
                             19870113
                                            US 1985-757763
                                                              19850722
     US 4636501
                       Α
                                         SE 1981-7574
                                                              19811217
PRIORITY APPLN. INFO.:
                                         EP 1982-850257
                                                              19821210
                                         US 1982-450006
                                                              19821215
                                         US 1983-482266
                                                              19830405
                                         US 1984-621147
                                                              19840618
AB
     Phenoxyisopropanolamines I [R = H, alkyl, cycloalkyl, cycloalkylalkyl; R1
     = H, Cl, Br, F; R2 = H, protective acyl group; R3, R4 = H, acyl,
     acyloxyalkyl; n = 2-4; R5, R6 = H, alkyl, hydroxyalkyl, alkoxyalkyl;
NR5R6
     = morpholino], useful as cardiovascular agents (no data), were prepd. by
     different methods. Glycidyl 4-(2-methoxyethoxy)phenyl ether was treated
     with N-(2-aminoethyl)-4-morpholinecarboxamide to give I (R-R4 = H, n = 2,
     NR5R6 = morpholino).
ΙT
     75930-29-9
     RL: RCT (Reactant)
        (ring cleavage by, of glycidyl aryl ethers)
RN
     75930-29-9 CAPLUS
     Urea, N-(2-aminoethyl)-N'-methyl- (9CI) (CA INDEX NAME)
CN
     0
'MeNH-C-NH-CH<sub>2</sub>-CH<sub>2</sub>-NH<sub>2</sub>
L23 ANSWER 8 OF 10 CAPLUS COPYRIGHT 2001 ACS
ACCESSION NUMBER:
                          1982:544557 CAPLUS
DOCUMENT NUMBER:
                          97:144557
TITLE:
                          3-Aminopropoxyphenyl derivatives and pharmaceutical
                          compositions containing them
INVENTOR(S):
                          Berthold, Richard; Louis, William John
                          Sandoz A.-G., Switz.
PATENT ASSIGNEE(S):
                          Eur. Pat. Appl., 57 pp.
SOURCE:
                          CODEN: EPXXDW
DOCUMENT TYPE:
                          Patent
                          English
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
```

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 52072	A1	19820519	EP 1981-810439	19811102
EP 52072	B1	19850220		
R: AT, BE,	CH, DE	, FR, GB,	IT, LU, NL, SE	
FI 8103412	À	19820507	FI 1981-3412	19811030
FI 76551	В	19880729		

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19881110
     FI 76551
                       С
     AT 11908
                       E
                            19850315
                                           AT 1981-810439
                                                             19811102
     DK 8104893
                       Α
                            19820507
                                           DK 1981-4893
                                                             19811104
                       В
                            19910624
     DK 161310
                       С
                            19911202
     DK 161310
     JP 57108047
                       A2
                            19820705
                                            JP 1981-175825
                                                             19811104
     JP 02057540
                       B4
                            19901205
                                           ES 1981-506842
     ES 506842
                       A1
                            19831201
                                                             19811104
                            19840110
                                           US 1981-318292
                                                             19811104
     US 4425362
                       Α
                                           IL 1981-64213
                                                             19811104
     IL 64213
                       Α1
                            19860731
     HU 28421
                       0
                            19831228
                                           HU 1981-3306
                                                             19811105
     HU 185976
                       В
                            19850428
     SU 1160933
                       A3
                            19850607
                                            SU 1981-3372598
                                                             19811105
                       Α1
                            19861104
                                            CA 1981-389517
                                                             19811105
     CA 1213594
     AU 8177171
                       A1
                            19820513
                                           AU 1981-77171
                                                             19811106
     AU 555619
                       B2
                            19861002
     ZA 8107702
                                            ZA 1981-7702
                                                             19811106
                       Α
                            19830629
     US 5347050
                            19940913
                                           US 1993-46937
                                                             19930413
                       Α
                                        CH 1980-8249
                                                             19801106
PRIORITY APPLN. INFO.:
                                        CH 1980-9347
                                                             19801218
                                        CH 1981-4073
                                                             19810619
                                        CH 1981-4074
                                                             19810619
                                        EP 1981-810439
                                                             19811102
                                        US 1981-318292
                                                             19811104
                                        US 1984-567471
                                                             19840103
                                        US 1985-778831
                                                             19850923
                                        US 1986-897557
                                                             19860818
                                        US 1988-173845
                                                             19880328
                                        US 1989-307028
                                                             19890203
                                        US 1989-399721
                                                             19890825
                                        US 1990-474185
                                                             19900202
                                        US 1990-584306
                                                             19900917
                                        US 1991-782791
                                                             19911021
     Title compds. I [R = alkyl, cycloalkyl, alkenyl, cycloalkylalkyl,
AΒ
     (un) substituted aryl, aralkyl, aralkenyl; R1 = H, substituent; R2 = H, R;
     Q = alkylene; Q1 = bond, NH; Y = O, S; Z = O, n = 2, 3; Z = bond, n =
     when R2 = cycloalkylalkyl, R = alkyl, and Q1 = bond, the R1 =
     or their physiol. acceptable hydrolyzable derivs. in esterified form, in
     either basic or salt forms, useful as cardioselective .beta .-
     adrenoreceptor blocking agents, were prepd. E.g., 4-PhCH2OC6H4OH was
     treated with 2-chloroethyl cyclopropylmethyl ether, and the resultant
     1-benzyloxy-4-(2-cyclopropylmethoxyethoxy) benzene was debenzylated by
     hydrogenolysis. The resulting 4-(2-cyclopropylmethoxyethoxy)phenol was
     brominated, benzylated and treated with CuCN to give 2-benzyloxy-5-(2-
     cyclopropylmethoxyethoxy) benzonitrile. The latter was debenzylated by
     hydrogenolysis and treated with epichlorohydrin to give
     2-(2,3-epoxypropoxy)-5-(2-cyclopropylmethoxyethoxy)benzonitrile.
     latter was fused with 1-(2-aminoethyl)-3-phenylurea to give
     1-[2-cyano-4-(2-cyclopropylmethoxyethoxy)phenoxy]-3-[2-(3-
     phenylureido) ethylamino] - 2-propanol (II). II was an effective
     .beta.-adrenoreceptor blocking agent. Ca. 82 examples of I were prepd.
     from the corresponding epoxy compds.
ΙT
```

RL: RCT (Reactant)

75930-29-9

```
(reaction of, with (epoxypropoxy)benzene derivs.)
RN
     75930-29-9 CAPLUS
     Urea, N-(2-aminoethyl)-N'-methyl- (9CI) (CA INDEX NAME)
CN
MeNH-C-NH-CH_2-CH_2-NH_2
L23 ANSWER 9 OF 10 CAPLUS COPYRIGHT 2001 ACS
                         1981:57984 CAPLUS
ACCESSION NUMBER:
DOCUMENT NUMBER:
                         94:57984
                         Potential inhibitors of nucleotide biosynthesis. 1.
TITLE:
                         Nitrosoureidonucleosides. 2
                         Montgomery, John A.; Thomas, H. Jeanette; Brockman,
AUTHOR(S):
                         Wallace; Wheeler, Glynn P.
                         Kettering-Meyer Lab., South. Res. Inst., Birmingham,
CORPORATE SOURCE:
                         AL, 35255, USA
                         J. Med. Chem. (1981), 24(2), 184-9
SOURCE:
                         CODEN: JMCMAR; ISSN: 0022-2623
DOCUMENT TYPE:
                         Journal
LANGUAGE:
                         English
     The title compds. I (R = H, Me, or cyclohexyl; R1 and R2 = H or N0; R3 = H
     hypoxanthin-9-yl, thymin-1-yl, or uracil-1-yl; R4 = H or OH) were prepd.
     and evaluated for alkylating activity. The low level of biol. activity
of
     I is apparently due to their stability compared to the known nitrosourea
     compds.
IT
     75930-29-9P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
        (prepn. and acylation of, by ribofuranuronic acid)
     75930-29-9 CAPLUS
RN
     Urea, N-(2-aminoethyl)-N'-methyl- (9CI) (CA INDEX NAME)
CN
MeNH-C-NH-CH2-CH2-NH2
IT
     75930-38-0P
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (prepn. of)
RN
     75930-38-0 CAPLUS
     Urea, N-(2-aminoethyl)-N'-methyl-, compd. with 2,4,6-trinitrophenol (1:1)
     (9CI) (CA INDEX NAME)
     CM
          1
     CRN 75930-29-9
     CMF C4 H11 N3 O
```

09/350,193

CM 2

CRN 88-89-1 CMF C6 H3 N3 O7

L23 ANSWER 10 OF 10 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

1978:590538 CAPLUS

DOCUMENT NUMBER:

89:190538

TITLE:

Method for the photometric determination of

N-monosubstituted carbamates

AUTHOR(S):

Schoene, K.; Steinhanses, J.

CORPORATE SOURCE:

Inst. Aerobiol., Fraunhofer-Ges., Schmallenberg, Ger.

SOURCE:

Fresenius' Z. Anal. Chem. (1978), 292(1), 29-33

CODEN: ZACFAU; ISSN: 0016-1152

DOCUMENT TYPE:

Journal

LANGUAGE: German

N-monosubstituted carbamates were converted to urea derivs. by reaction with 1,3-diaminopropane in the presence of small amts. of NaOH. The urea derivs. were detd. spectrophotometrically at 577 nm by using the carbamide

reaction described by W. R. Fearon (1939). The detection limit for N-methylcarbamates is in the range of 20 nmols. In the case of N-methylcarbamates, N-methyl-N'-(3-aminopropyl)urea was found to be the intermediate urea deriv., which is formed in nearly quant. yield. The amidation reaction mechanism of the N-methylcarbamates was studied on N-methylurethane.

68156-37-6P IT

> RL: ANST (Analytical study); PREP (Preparation) (prepn. of)

68156-37-6 CAPLUS RN

Urea, N-(3-aminopropyl)-N'-methyl- (9CI) (CA INDEX NAME) CN

=> d his (FILE 'HOME' ENTERED AT 10:38:39 ON 01 JUN 2001) FILE 'REGISTRY' ENTERED AT 10:38:44 ON 01 JUN 2001 STRUCTURE UPLOADED L1L2 50 S L1 36270 S L1 FULL L3 L4STRUCTURE UPLOADED L5 14060 S L4 FULL SUB=L3 L6 5399 S L5 AND 3/N L7 734 S ETHYL(L) DIMETHYL(L) AMINO(L) PROPYL(L) UREA 0 S UREA, "N-ETHYL-N'-(3-DIMETHYLAMINOPROPYL)-" L8 L9 4 S ETHYL(L) DIMETHYLAMINOPROPYL(L) UREA L10 3 S L9 AND 1/NC L11 1 S L10 AND 1/0 FILE 'CAPLUS' ENTERED AT 10:52:57 ON 01 JUN 2001 L12 15 S L11 L13 2 S L11/THU FILE 'USPATFULL' ENTERED AT 10:55:19 ON 01 JUN 2001 L14 2 S L11 FILE 'MARPAT' ENTERED AT 10:56:49 ON 01 JUN 2001 L15 0 S L11 L16 0 S L9 FILE 'BEILSTEIN' ENTERED AT 10:57:15 ON 01 JUN 2001 L17 1 S L11 FILE 'REGISTRY' ENTERED AT 10:57:49 ON 01 JUN 2001 STRUCTURE UPLOADED L18 L19 11247 S L18 FULL SUB=L3 L2050 S L18 L21 0 S L18 CSS 8 S L18 CSS FULL L22 FILE 'CAPLUS' ENTERED AT 11:01:43 ON 01 JUN 2001 · L23 10 S L22 => s 122/thu 10 L22 375285 THU/RL 0 L22/THU L24 (L22 (L) THU/RL) => file uspatful COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION FULL ESTIMATED COST 44.43 598.50

SINCE FILE

ENTRY

-5.88

TOTAL

SESSION

-15.88

DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)

CA SUBSCRIBER PRICE

FILE 'USPATFULL' ENTERED AT 11:03:35 ON 01 JUN 2001 CA INDEXING COPYRIGHT (C) 2001 AMERICAN CHEMICAL SOCIETY (ACS)

FILE COVERS 1971 TO PATENT PUBLICATION DATE: 29 May 2001 (20010529/PD) FILE LAST UPDATED: 29 May 2001 (20010529/ED) HIGHEST PATENT NUMBER: US8411134 CA INDEXING IS CURRENT THROUGH 29 May 2001 (20010529/UPCA) ISSUE CLASS FIELDS (/INCL) CURRENT THROUGH: 29 May 2001 (20010529/PD) REVISED CLASS FIELDS (/NCL) LAST RELOADED: Apr 2001 USPTO MANUAL OF CLASSIFICATIONS THESAURUS ISSUE DATE: Apr 2001

>>> Page images are available for patents from 1/1/1997. Current <<< >>> week patent text is typically loaded by Thursday morning and <<< >>> page images are available for display by the end of the day. <<< >>> Image data for the /FA field are available the following week. <<<

>>> Complete CA file indexing for chemical patents (or equivalents) <<< >>> is included in file records. A thesaurus is available for the <<<>>> USPTO Manual of Classifications in the /NCL, /INCL, and /RPCL <<< >>> fields. This thesaurus includes catchword terms from the <<< >>> USPTO/MOC subject headings and subheadings. Thesauri are also <<< >>> available for the WIPO International Patent Classification <<< >>> (IPC) Manuals, editions 1-6, in the /IC1, /IC2, /IC3, /IC4, <<< >>> /IC5, and /IC (/IC6) fields, respectively. The thesauri in <<< >>> the /IC5 and /IC fields include the corresponding catchword <<< >>> terms from the IPC subject headings and subheadings. <<<

This file contains CAS Registry Numbers for easy and accurate substance identification.

=> s 122 L25

7 L22

=> d ibib ab hitstr 1-7

L25 ANSWER 1 OF 7 USPATFULL

ACCESSION NUMBER: 94:80134 USPATFULL

TITLE: 3-aminopropoxyphen

and

3-aminopropoxyphenyl derivatives, their preparation

pharmaceutical compositions containing them

INVENTOR(S): Berthold, Richard, 9 Ahornstrasse, CH-4103 Bottmingen,

Switzerland

Louis, William J., 3 Balmoral Avenue, Kew, 3101

Victoria, Australia

NUMBER DATE

PATENT INFORMATION:
APPLICATION INFO.:
RELATED APPLN. INFO.:

US 5347050 19940913 US 1993-46937 19930413 (8)

Continuation of Ser. No. US 1991-782791, filed on 21 Oct 1991, now abandoned which is a continuation of

Ser.

No. US 1990-584306, filed on 17 Sep 1990, now

abandoned

which is a continuation of Ser. No. US 1990-474185, filed on 2 Feb 1990, now abandoned which is a

continuation of Ser. No. US 1989-399721, filed on 25 Aug 1989, now abandoned which is a continuation of

Ser.

No. US 1989-307028, filed on 3 Feb 1989, now abandoned which is a continuation of Ser. No. US 1988-173845, filed on 28 Mar 1988, now abandoned which is a continuation of Ser. No. US 1986-897557, filed on 18 Aug 1986, now abandoned which is a continuation of

Ser.

No. US 1985-778831, filed on 23 Sep 1985, now

abandoned

which is a continuation of Ser. No. US 1984-567471, filed on 3 Jan 1984, now abandoned which is a division of Ser. No. US 1981-318292, filed on 4 Nov 1981, now patented, Pat. No. US 4425362

	NUMBER	DATE
PRIORITY INFORMATION:	CH 1980-8249	19801106
	CH 1980-9347	19801218
	CH 1981-4073	19810619
	CH 1991-407481	19910619
DOCUMENT TYPE:	Utility	
PRIMARY EXAMINER:	Dees, Jose G.	

PRIMARY EXAMINER: Dees, Jose G.
ASSISTANT EXAMINER: Carr, Deborah D.
LEGAL DEPORTSENTATIVE: Sughana Micro 7

LEGAL REPRESENTATIVE: Sughrue, Mion, Zinn, Macpeak & Seas

NUMBER OF CLAIMS: 7
EXEMPLARY CLAIM: 1
LINE COUNT: 1090

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The compounds of formula I, ##STR1## wherein the substituents have various significances, and physiologically acceptable hydrolyzable derivatives thereof having the hydroxy group in the 2 position of the 3-aminopropoxy side chain in esterified form, are useful as cardioselective .beta.-adrenoceptor blocking agents.

IT 75930-29-9

(reaction of, with (epoxypropoxy)benzene derivs.)

RN 75930-29-9 USPATFULL

CN Urea, N-(2-aminoethyl)-N'-methyl- (9CI) (CA INDEX NAME)

 $\begin{tabular}{l} \tt O \\ || \\ \tt MeNH-C-NH-CH_2-CH_2-NH_2 \\ \end{tabular}$

L25 ANSWER 2 OF 7 USPATFULL

ACCESSION NUMBER: 90:48806 USPATFULL

TITLE: 2-hydroxypropylamine aryl ester derivatives and

pharmaceutical use

INVENTOR(S): Kam, Sheung T., Vernon Hills, IL, United States

Matier, William L., Libertyville, IL, United States E. I. Du Pont de Nemours and Company, Wilmington, DE,

PATENT ASSIGNEE(S): E. I. Du Pont de Nemours and Compunited States (U.S. corporation)

RELATED APPLN. INFO.:

NUMBER DATE US 4935421 19900619 PATENT INFORMATION: US 1989-318147 19890301 (7) APPLICATION INFO.: Division of Ser. No. US 1986-838082, filed on 10 Mar RELATED APPLN. INFO.: 1986, now patented, Pat. No. US 4810717 which is a division of Ser. No. US 1981-320773, filed on 21 Nov 1981, now patented, Pat. No. US 4582855 DOCUMENT TYPE: Utility PRIMARY EXAMINER: Ramsuer, Robert W. NUMBER OF CLAIMS: 27 EXEMPLARY CLAIM: 1 LINE COUNT: 1470 CAS INDEXING IS AVAILABLE FOR THIS PATENT. Novel compounds of the general formula ##STR1## wherein Ar represents a substituted or unsubstituted aromatic or heterocyclic group; W represents alkylene of from 1 to about 10 carbon atoms; and B represents --NR.sub.2 COR.sub.1, --NR.sub.2 CONR.sub.1 R.sub.3, --NR.sub.2 SO.sub.2 R.sub.1, --NR.sub.2 SO.sub.2 NR.sub.1 R.sub.3, or --NR.sub.2 COOR.sub.1 wherein R.sub.1, R.sub.2 and R.sub.3 may be the same or different and may be hydrogen, alkyl, alkoxyalkyl, cycloalkyl, alkenyl, alkynyl, aryl, heteroaryl, or aralkyl, except that R.sub.1 is not hydrogen when B is --NR.sub.2 SO.sub.2 R.sub.1 or --NR.sub.2 COOR.sub.1, or R.sub.1 and R.sub.3 may together with N form a 5 to 7 membered heterocyclic group; and the pharmaceutically acceptable salts thereof. These compounds exhibit .beta.-adrenergic blocking activity and are also useful in the treatment of glaucoma. IT 75930-29-9P (prepn. of) RN 75930-29-9 USPATFULL Urea, N-(2-aminoethyl)-N'-methyl- (9CI) (CA INDEX NAME) CN MeNH-C-NH-CH2-CH2-NH2 L25 ANSWER 3 OF 7 USPATFULL ACCESSION NUMBER: 89:17318 USPATFULL TITLE: 2-hydroxypropylamine aryl ester derivatives INVENTOR(S): Kam, Sheung T., Vernon Hills, IL, United States Matier, William L., Libertyville, IL, United States PATENT ASSIGNEE(S): E. I. du Pont de Nemours and Company, Wilmington, DE, United States (U.S. corporation) NUMBER US 4810717 19890307 US 1986-838082 19860310 (6) Division of Ser. No. US 1981-320773, filed on 12 Nov PATENT INFORMATION: APPLICATION INFO.:

1981, now patented, Pat. No. US 4582855

DOCUMENT TYPE:

Utility

PRIMARY EXAMINER:

Lee, Mary C.

ASSISTANT EXAMINER:

Whittenbaugh, Robert C.

LEGAL REPRESENTATIVE:

Fato, Gildo E.

NUMBER OF CLAIMS:

33 1,11

EXEMPLARY CLAIM:

LINE COUNT:

1764

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Novel compounds of the general formula ##STR1## wherein Ar represents a substituted or unsubstituted aromatic or heterocyclic group; W represents alkylene of from 1 to about 10 carbon atoms; and B

represents

--NR.sub.2 COR.sub.1, --NR.sub.2 CONR.sub.1 R.sub.3, --NR.sub.2

SO.sub.2

R.sub.1, --NR.sub.2 SO.sub.2 NR.sub.1 R.sub.3, or --NR.sub.2 COOR.sub.1 wherein R.sub.1, R.sub.2 and R.sub.3 may be the same or different and may be hydrogen, alkyl, alkoxyalkyl, cycloalkyl, alkenyl, alkynyl,

aryl,

heteroaryl, or aralkyl, except that R.sub.1 is not hydrogen when B is --NR.sub.2 SO.sub.2 R.sub.1 or --NR.sub.2 COOR.sub.1, or R.sub.1 and R.sub.3 may together with N form a 5 to 7 membered heterocyclic group; and the pharmaceutically acceptable salts thereof. These compounds exhibit .beta.-adrenergic blocking activity and are also useful in the treatment of glaucoma.

75930-29-9P

(prepn. of)

RN 75930-29-9 USPATFULL

CN Urea, N-(2-aminoethyl)-N'-methyl- (9CI) (CA INDEX NAME)

0 MeNH-C-NH-CH2-CH2-NH2

L25 ANSWER 4 OF 7 USPATFULL

ACCESSION NUMBER:

89:4612 USPATFULL

TITLE: INVENTOR(S): 2-hydroxypropylamine heteroaryl ester derivatives

Kam, Sheung T., Chicago, IL, United States

Matier, William L., Libertyville, IL, United States Patil, Ghanshyam, Vernon Hills, IL, United States Mai, Khuong H. X., Waukegan, IL, United States

PATENT ASSIGNEE(S):

E. I. Du Pont de Nemours and Company, Wilmington, DE,

United States (U.S. corporation)

NUMBER

PATENT INFORMATION:

US 4798892

19890117

APPLICATION INFO.:

US 1986-851629 19860414 (6)

RELATED APPLN. INFO.:

Continuation-in-part of Ser. No. US 1981-320773, filed on 12 Nov 1981, now patented, Pat. No. US 4582855,

issued on 15 Apr 1986

DOCUMENT TYPE:

Utility

PRIMARY EXAMINER:

Raymond, Richard L.

LEGAL REPRESENTATIVE:

Fato, Gildo E.

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

1

LINE COUNT:

1391

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention relates to compounds of the general formula ##STR1## wherein Ar represents a substituted or unsubstituted heterocyclic group; W represents alkylene of from 1 to about 10 carbon atoms; and B represents --NR.sub.2 COR.sub.1, --NR.sub.2 CONR.sub.1 R.sub.3, --NR.sub.2 SO.sub.2 R.sub.1, NR.sub.2 SO.sub.2 NR.sub.1 R.sub.3, or --NR.sub.2 COOR.sub.1, wherein R.sub.1, R.sub.2 and R.sub.3 may be alike or different and may be hydrogen, alkyl, alkoxyalkyl cycloalkyl, alkenyl, alkynyl, aryl, heteroaryl, or aralkyl, except that R.sub.1 is not hydrogen when B is --NR.sub.2 SO.sub.2 R.sub.1 or --NR.sub.2 COOR.sub.1, or R.sub.1 and R.sub.3 may together with N form

а

5 to 7 membered heterocyclic group and the pharmaceutically acceptable salts thereof. The compounds exhibit beta-adrenergic blocking activity and are also useful in the treatment of glaucoma.

122036-80-0P

(prepn. and reaction of, in prepn. of .beta.-adrenergic blockers)

RN 122036-80-0 USPATFULL

Urea, N-(2-aminoethyl)-N'-methyl-, monohydrochloride (9CI) (CA INDEX

MeNH-C-NH-CH2-CH2-NH2

HCl

L25 ANSWER 5 OF 7 USPATFULL

ACCESSION NUMBER:

87:3253 USPATFULL

TITLE:

Para-substituted 3-phenoxy-1-carbonylamino-alkylamino-

propanol compounds, beta receptor blocking

compositions

INVENTOR(S):

Gustafsson, Bill B. R., Bollebygd, Sweden

Hedberg, Sven A., Gr.ang.bo, Sweden Lundgren, Bo T., Frilles.ang.s, Sweden

PATENT ASSIGNEE(S):

Aktiebolaget Hassle, Molndal, Sweden (non-U.S.

corporation)

NUMBER DATE

PATENT INFORMATION:

US 4636501

19870113

APPLICATION INFO.: RELATED APPLN. INFO.: US 1985-757763 19850722 (6)

Continuation of Ser. No. US 1984-621147, filed on 18 Jun 1984, now abandoned which is a continuation of

Ser.

No. US 1983-482266, filed on 5 Apr 1983, now abandoned

which is a continuation-in-part of Ser. No. US

1982-450006, filed on 15 Dec 1982, now abandoned

NUMBER DATE PRIORITY INFORMATION: SE 1981-7574 19811217 DOCUMENT TYPE: Utility PRIMARY EXAMINER: Ramsuer, Robert W. Brumbaugh, Graves, Donohue & Raymond LEGAL REPRESENTATIVE: NUMBER OF CLAIMS: EXEMPLARY CLAIM: 1.9 LINE COUNT: 1017 CAS INDEXING IS AVAILABLE FOR THIS PATENT. Compounds of the formula ##STR1## having beta receptor blocking properties, are disclosed. 75930-29-9 (ring cleavage by, of glycidyl aryl ethers) RN 75930-29-9 USPATFULL CN Urea, N-(2-aminoethyl)-N'-methyl- (9CI) (CA INDEX NAME) 0 $MeNH-C-NH-CH_2-CH_2-NH_2$ L25 ANSWER 6 OF 7 USPATFULL ACCESSION NUMBER: 86:21877 USPATFULL TITLE: Aromatic and esters of hydroxypropylamines INVENTOR(S): Kam, Sheung T., Vernon Hills, IL, United States Matier, William L., Libertyville, IL, United States American Hospital Supply Corporation, Evanston, IL, PATENT ASSIGNEE(S): United States (U.S. corporation) NUMBER DATE PATENT INFORMATION: US 4582855 19860415 US 1981-320773 19811112 (6) APPLICATION INFO.: DOCUMENT TYPE: Utility PRIMARY EXAMINER: Jiles, Henry R. ASSISTANT EXAMINER: Whittenbaugh, Robert C. LEGAL REPRESENTATIVE: Kanady, Mary Jo; Barbeau, Donald L.; Fato, Gildo E. NUMBER OF CLAIMS: EXEMPLARY CLAIM: 1,30 LINE COUNT: 1804 CAS INDEXING IS AVAILABLE FOR THIS PATENT. Novel compounds of the general formula ##STR1## wherein Ar represents a substituted or unsubstituted aromatic or heterocyclic group; W represents alkylene of from 1 to about 10 carbon atoms; and B --NR.sub.2 COR.sub.1, --NR.sub.2 CONR.sub.1 R.sub.3, --NR.sub.2 SO.sub.2 R.sub.1, --NR.sub.2 SO.sub.2 NR.sub.1 R.sub.3, or --NR.sub.2 COOR.sub.1 wherein R.sub.1, R.sub.2 and R.sub.3 may be the same or different and may be hydrogen, alkyl, alkoxyalkyl, cycloalkyl, alkenyl, alkynyl, aryl,

heteroaryl, or aralkyl, except that R.sub.1 is not hydrogen when B is --NR.sub.2 SO.sub.2 R.sub.1 or --NR.sub.2 COOR.sub.1, or R.sub.1 and R.sub.3 may together with N form a 5 to 7 momoered heterocyclic group; and the pharmaceutically acceptable salts thereof. These compounds exhibit .beta.-adrenergic blocking activity and are also useful in the treatment of glaucoma.

IT 75930-29-9P

(prepn. of)

75930-29-9 USPATFULL

Urea, N-(2-aminoethyl)-N'-methyl- (9CI) (CA INDEX NAME)

0 MeNH-C-NH-CH2-CH2-NH2

L25 ANSWER 7 OF 7 USPATFULL

ACCESSION NUMBER: 84:2057 USPATFULL

TITLE:

3-Aminopropoxyphenyl derivatives and pharmaceutical

compositions containing them

INVENTOR(S): Berthold, Richard, Bottmingen, Switzerland

Louis, William J., Kew, Australia

PATENT ASSIGNEE(S): Sandoz Ltd., Basel, Switzerland (non-U.S. corporation)

NUMBER DATE ______ PATENT INFORMATION: US 4425362 19840110 APPLICATION INFO.: US 1981-318292 19811104 (6)

NUMBER DATE ______ CH 1980-8249 19801106 CH 1980-9347 19801218 CH 1981-4073 19810619 PRIORITY INFORMATION: 19801218 19810619 19810619 CH 1981-4074 DOCUMENT TYPE:

Utility

PRIMARY EXAMINER: Torrence, Dolph H.

LEGAL REPRESENTATIVE: Sharkin, Gerald D.; Honor, Robert S.

NUMBER OF CLAIMS: EXEMPLARY CLAIM: 1,10 LINE COUNT: 1101

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The compounds of formula I, ##STR1## wherein the substituents have various significances, and physiologically acceptable hydrolyzable derivatives thereof having the hydroxy group in the 2 position of the 3-aminopropoxy side chain in esterified form, are useful as cardioselective .beta.-adrenoceptor blocking agents.

75930-29-9

(reaction of, with (epoxypropoxy)benzene derivs.)

RN 75930-29-9 USPATFULL

Urea, N-(2-aminoethyl)-N'-methyl- (9CI) (CA INDEX NAME) CN

=> file reg COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION FULL ESTIMATED COST 36.50 635.00 DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) SINCE FILE TOTAL ENTRY SESSION CA SUBSCRIBER PRICE 0.00 -15.88

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=> d his

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FILE 'USPATFULL' ENTERED AT 10:55:19 ON 01 JUN 2001 L14 2 S L11

FILE 'MARPAT' ENTERED AT 10:56:49 ON 01 JUN 2001 L15 0 S L11 L16 0 S L9 FILE 'BEILSTEIN' ENTERED AT 10:57:15 ON 01 JUN 2001 L17 FILE 'REGISTRY' ENTERED AT 10:57:49 ON 01 JUN 2001 L18 STRUCTURE UPLOADED 11247 S L18 FULL SUB=L3 L19 L20 50 S L18 L21 0 S L18 CSS L22 8 S L18 CSS FULL FILE 'CAPLUS' ENTERED AT 11:01:43 ON 01 JUN 2001 L23 10 S L22 L24 0 S L22/THU FILE 'USPATFULL' ENTERED AT 11:03:35 ON 01 JUN 2001 L25 7 S L22 FILE 'REGISTRY' ENTERED AT 11:05:41 ON 01 JUN 2001 => s 118 full FULL SEARCH INITIATED 11:05:52 FILE 'REGISTRY' FULL SCREEN SEARCH COMPLETED - 54245 TO ITERATE 100.0% PROCESSED 54245 ITERATIONS 11247 ANSWERS SEARCH TIME: 00.00.05 L26 11247 SEA SSS FUL L18 => d scan L26 11247 ANSWERS REGISTRY COPYRIGHT 2001 ACS Urea, N-[1-[(3,4-dichlorophenyl)methyl]-1,6-dihydro-6-oxo-3-pyridinyl]-N'-1[3-(dimethylamino)propyl]- (9CI) MF C18 H22 C12 N4 O2

$$Me_2N - (CH_2)_3 - NH - C - NH$$
 $N - CH_2$
 $C1$

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):2

L26 11247 ANSWERS REGISTRY COPYRIGHT 2001 ACS
IN Urea, N-[2-[[[(4-methylphenyl)amino]carbonyl]amino]ethyl]-N'-(4-

nitrophenyl)-N-(4-phenoxyphenyl)- (9CI) MF C29 H27 N5 O5

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

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L27 STRUCTURE UPLOADED

=> d his

(FILE 'HOME' ENTERED AT 10:38:39 ON 01 JUN 2001)

FILE 'REGISTRY' ENTERED AT 10:38:44 ON 01 JUN 2001 L1STRUCTURE UPLOADED L2 50 S L1 L336270 S L1 FULL L4STRUCTURE UPLOADED L514060 S L4 FULL SUB=L3 L6 5399 S L5 AND 3/N L7 734 S ETHYL(L) DIMETHYL(L) AMINO(L) PROPYL(L) UREA $rac{1}{8}$ 0 S UREA, "N-ETHYL-N'-(3-DIMETHYLAMINOPROPYL)-" L9 4 S ETHYL(L) DIMETHYLAMINOPROPYL(L) UREA

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3 S L9 AND 1/NC
1 S L10 AND 1/O
L10
L11
     FILE 'CAPLUS' ENTERED AT 10:52:57 ON 01 JUN 2001
L12
             15 S L11
L13
             2 S L11/THU
     FILE 'USPATFULL' ENTERED AT 10:55:19 ON 01 JUN 2001
L14
              2 S L11
     FILE 'MARPAT' ENTERED AT 10:56:49 ON 01 JUN 2001
L15
              0 S L11
L16
              0 S L9
     FILE 'BEILSTEIN' ENTERED AT 10:57:15 ON 01 JUN 2001
L17
              1 S L11
     FILE 'REGISTRY' ENTERED AT 10:57:49 ON 01 JUN 2001
               STRUCTURE UPLOADED
L19
          11247 S L18 FULL SUB=L3
L20
            50 S L18
L21
             0 S L18 CSS
L22
              8 S L18 CSS FULL
     FILE 'CAPLUS' ENTERED AT 11:01:43 ON 01 JUN 2001
L23
           10 S L22
L24
             0 S L22/THU
    FILE 'USPATFULL' ENTERED AT 11:03:35 ON 01 JUN 2001
L25
              7 S L22
    FILE 'REGISTRY' ENTERED AT 11:05:41 ON 01 JUN 2001
L26
     11247 S L18 FULL
               STRUCTURE UPLOADED
L27
=> s 127 sub=126 full
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FULL SUBSET SCREEN SEARCH COMPLETED - 11247 TO ITERATE
100.0% PROCESSED 11247 ITERATIONS
                                                            3145 ANSWERS
SEARCH TIME: 00.00.02
         3145 SEA SUB=L26 SSS FUL L27
=> d scan
L28 3145 ANSWERS REGISTRY COPYRIGHT 2001 ACS
IN Urea,
N-[6-bromo-2-(3-thienyl)-4-quinolinyl]-N'-[3-(dimethylamino)propyl]-
MF C19 H21 Br N4 O S
```

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):1

L28 3145 ANSWERS REGISTRY COPYRIGHT 2001 ACS
IN Urea, N-(2,3-dichlorophenyl)-N'-[6-[(4-nitrophenyl)amino]hexyl]- (9CI)
MF C19 H22 C12 N4 O3

$$\begin{array}{c|c} & & & & & & & & & & \\ & & & & & & & \\ & & & & & & \\ & & & & & & \\ & & & & & & \\ & & & & & & \\ & & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & \\ & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ &$$

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

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L14

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L29 STRUCTURE UPLOADED

2 S L11

=> d his

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FILE 'REGISTRY' ENTERED AT 10:38:44 ON 01 JUN 2001 L1STRUCTURE UPLOADED L2 50 S L1 L3 36270 S L1 FULL L4STRUCTURE UPLOADED L5 14060 S L4 FULL SUB=L3 L6 5399 S L5 AND 3/N L7 734 S ETHYL(L) DIMETHYL(L) AMINO(L) PROPYL(L) UREA 0 S UREA, "N-ETHYL-N'-(3-DIMETHYLAMINOPROPYL)-" L8 4 S ETHYL(L)DIMETHYLAMINOPROPYL(L)UREA L9 L10 3 S L9 AND 1/NC L11 1 S L10 AND 1/0 FILE 'CAPLUS' ENTERED AT 10:52:57 ON 01 JUN 2001 L12 15 S L11 L13 2 S L11/THU FILE 'USPATFULL' ENTERED AT 10:55:19 ON 01 JUN 2001

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FILE 'MARPAT' ENTERED AT 10:56:49 ON 01 JUN 2001
L15
              0 S L11
L16
              0 S L9
     FILE 'BEILSTEIN' ENTERED AT 10:57:15 ON 01 JUN 2001
L17
              1 S L11
     FILE 'REGISTRY' ENTERED AT 10:57:49 ON 01 JUN 2001
L18
                STRUCTURE UPLOADED
          11247 S L18 FULL SUB=L3
L19
L20
             50 S L18
L21
              0 S L18 CSS
L22
              8 S L18 CSS FULL
     FILE 'CAPLUS' ENTERED AT 11:01:43 ON 01 JUN 2001
L23
             10 S L22
L24
              0 S L22/THU
     FILE 'USPATFULL' ENTERED AT 11:03:35 ON 01 JUN 2001
L25
              7 S L22
     FILE 'REGISTRY' ENTERED AT 11:05:41 ON 01 JUN 2001
L26
          11247 S L18 FULL
L27
                STRUCTURE UPLOADED
L28
           3145 S L27 FULL SUB=L26
L29
                STRUCTURE UPLOADED
=> s 129 sub=128 full
FULL SUBSET SEARCH INITIATED 11:13:36 FILE 'REGISTRY'
FULL SUBSET SCREEN SEARCH COMPLETED - 3145 TO ITERATE
100.0% PROCESSED
                    3145 ITERATIONS
                                                             1523 ANSWERS
SEARCH TIME: 00.00.04
L30
           1523 SEA SUB=L28 SSS FUL L29
=> d scan
L30 1523 ANSWERS
                    REGISTRY COPYRIGHT 2001 ACS
     4,6,15,17-Tetraazaeicosane-1,3,18,20-tetracarboxylic acid, 5,16-dioxo-,
     3,18-dimethyl ester, (3S,18S)- (9CI)
     C22 H38 N4 O10
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Absolute stereochemistry.

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

=> d his (FILE 'HOME' ENTERED AT 10:38:39 ON 01 JUN 2001) FILE 'REGISTRY' ENTERED AT 10:38:44 ON 01 JUN 2001 L1 STRUCTURE UPLOADED L2 50 S L1 L3 36270 S L1 FULL L4STRUCTURE UPLOADED L5 14060 S L4 FULL SUB=L3 L6 5399 S L5 AND 3/N 734 S ETHYL(L) DIMETHYL(L) AMINO(L) PROPYL(L) UREA L7 0 S UREA, "N-ETHYL-N'-(3-DIMETHYLAMINOPROPYL)-" L8 L9 4 S ETHYL(L)DIMETHYLAMINOPROPYL(L)UREA L10 3 S L9 AND 1/NC L11 1 S L10 AND 1/O FILE 'CAPLUS' ENTERED AT 10:52:57 ON 01 JUN 2001 L12 15 S L11 L13 2 S L11/THU FILE 'USPATFULL' ENTERED AT 10:55:19 ON 01 JUN 2001 L14 2 S L11 FILE 'MARPAT' ENTERED AT 10:56:49 ON 01 JUN 2001 L15 0 S L11 L16 0 S L9 FILE 'BEILSTEIN' ENTERED AT 10:57:15 ON 01 JUN 2001 L17 1 S L11 FILE 'REGISTRY' ENTERED AT 10:57:49 ON 01 JUN 2001 L18 STRUCTURE UPLOADED 11247 S L18 FULL SUB=L3 L19 L20 50 S L18 L21 0 S L18 CSS L22 8 S L18 CSS FULL FILE 'CAPLUS' ENTERED AT 11:01:43 ON 01 JUN 2001 L23 10 S L22 L24 0 S L22/THU FILE 'USPATFULL' ENTERED AT 11:03:35 ON 01 JUN 2001 L25 7 S L22 FILE 'REGISTRY' ENTERED AT 11:05:41 ON 01 JUN 2001 L26 11247 S L18 FULL L27 STRUCTURE UPLOADED L28 3145 S L27 FULL SUB=L26 L29 STRUCTURE UPLOADED L30 1523 S L29 FULL SUB=L28

=> s 130 and 1/o2846307 1/0 L31 381 L30 AND 1/O

09/350,193

=> d scan

L31 381 ANSWERS REGISTRY COPYRIGHT 2001 ACS
IN 1-Dodecanaminium, N,N-dimethyl-N-[2-[[[[3-[(3,3,4,4,5,5,6,6,6-nonafluorohexyl)thio]propyl]amino]carbonyl]amino]ethyl]-, bromide (9CI)
MF C26 H47 F9 N3 O S . Br

● Br-

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

=> s 131 not s/els 4193743 S/ELS L32 331 L31 NOT S/ELS

=> d scan

L32 331 ANSWERS REGISTRY COPYRIGHT 2001 ACS
IN Urea, N-(4-aminobutyl)-N'-(triphenylmethyl)- (9CI)
MF C24 H27 N3 O

$$\begin{array}{c} \text{O} \\ || \\ \text{Ph}_{3}\text{C-NH-C-NH-(CH}_{2})_{4}-\text{NH}_{2} \end{array}$$

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):1

L32 331 ANSWERS REGISTRY COPYRIGHT 2001 ACS
IN Urea, N-[2-(dimethylamino)ethyl]-N'-ethyl-, monohydrochloride (9CI)
MF C7 H17 N3 O . Cl H

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):1

L32 331 ANSWERS REGISTRY COPYRIGHT 2001 ACS

IN 1-Butanaminium, N,N,N-trimethyl-4-[[(methylamino)carbonyl]amino]- (9CI)

MF C9 H22 N3 O

CI COM

O || MeNH-C-NH-(CH₂)4-N+Me₃

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

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FILE 'REGISTRY' ENTERED AT 10:38:44 ON 01 JUN 2001
L1
                STRUCTURE UPLOADED
L2
             50 S L1
L3
          36270 S L1 FULL
L4
                STRUCTURE UPLOADED
L5
          14060 S L4 FULL SUB=L3
L6
           5399 S L5 AND 3/N
L7
            734 S ETHYL(L) DIMETHYL(L) AMINO(L) PROPYL(L) UREA
L8
              0 S UREA, "N-ETHYL-N'-(3-DIMETHYLAMINOPROPYL)-"
L9
              4 S ETHYL(L) DIMETHYLAMINOPROPYL(L) UREA
L10
              3 S L9 AND 1/NC
L11
              1 S L10 AND 1/0
     FILE 'CAPLUS' ENTERED AT 10:52:57 ON 01 JUN 2001
L12
             15 S L11
L13
              2 S L11/THU
     FILE 'USPATFULL' ENTERED AT 10:55:19 ON 01 JUN 2001
L14
              2 S L11
     FILE 'MARPAT' ENTERED AT 10:56:49 ON 01 JUN 2001
L15
              0 S L11
L16
              0 S L9
     FILE 'BEILSTEIN' ENTERED AT 10:57:15 ON 01 JUN 2001
L17
              1 S L11
     FILE 'REGISTRY' ENTERED AT 10:57:49 ON 01 JUN 2001
L18
               STRUCTURE UPLOADED
L19
          11247 S L18 FULL SUB=L3
L20
             50 S L18
L21
              0 S L18 CSS
L22
              8 S L18 CSS FULL
     FILE 'CAPLUS' ENTERED AT 11:01:43 ON 01 JUN 2001
L23
             10 S L22
L24
              0 S L22/THU
     FILE 'USPATFULL' ENTERED AT 11:03:35 ON 01 JUN 2001
L25
              7 S L22
     FILE 'REGISTRY' ENTERED AT 11:05:41 ON 01 JUN 2001
L26
          11247 S L18 FULL
L27
                STRUCTURE UPLOADED
L28
           3145 S L27 FULL SUB=L26
L29
                STRUCTURE UPLOADED
L30
           1523 S L29 FULL SUB=L28
L31
           381 S L30 AND 1/0
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L32 331 S L31 NOT S/ELS

FILE 'CAPLUS' ENTERED AT 11:14:57 ON 01 JUN 2001

 \Rightarrow s 132/thu

226 L32

375285 THU/RL

L33

14 L32/THU

(L32 (L) THU/RL)

=> d ibib ab hitstr 1-14

L33 ANSWER 1 OF 14 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

2001:338479 CAPLUS

TITLE:

Preparation of amides and ureas as activators of

soluble quanylate cyclase

INVENTOR(S):

Selwood, David; Glen, Robert; Reynolds, Karen;

Wishart, Grant

PATENT ASSIGNEE(S):

University College London, UK

SOURCE:

PCT Int. Appl., 101 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

```
PATENT NO.
                                                APPLICATION NO. DATE
                       KIND DATE
      -----
     WO 2001032604
                         A1
                               20010510
                                                WO 2000-GB4249 20001106
          W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
              CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU,
              SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN,
              YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
          RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
              DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,
              BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
PRIORITY APPLN. INFO.:
                                             GB 1999-26286
                                                                 A 19991105
                                             US 2000-201382
                                                                 P 20000502
```

The title compds. R4PZNR1R2 [I; R1, R2 = alkyl; R1R2 together form AB alkylene; Z = alkylene; P = a direct bond, X, Y, W, XY, YW, XYW (wherein

= O, S, NR3; R3 = H, alkyl; Y = UV; V = a direct bond, alkylene; U = CS, CO, SO2, C(:NR); R = H, OH, alkyl; X = O, NR6; R6 = H, alkyl, alkenyl, etc.); R4 = alkyl, alkenyl, alkynyl, etc.], useful in the activation of sol. guanylate cyclase, were prepd. E.g., synthesis of the urea II, starting with 4-bromoaniline and 1-(3-aminopropyl)pyrrolidine, was given. Biol. data for compds. I (e.g., IC50 for inhibition of platelet aggregation) were presented.

IT 32897-26-0P 338980-63-5P

> RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of amides and ureas as activators of sol. guanylate cyclase) 32897-26-0 CAPLUS RN

CN Urea, N-[3-(dimethylamino)propyl]-N'-ethyl- (9CI) (CA INDEX NAME)

EtNH-C-NH-(CH₂)₃-NMe₂

RN338980-63-5 CAPLUS

Urea, N-[3-(dimethylamino)propyl]-N'-(1-pyrenylmethyl)- (9CI) (CA INDEX CN NAME)

REFERENCE COUNT:

24

REFERENCE(S):

1986,

(8) Catanese, B; BOLLETTINO CHIMICO FARMACEUTICO

V125(7), P228 CAPLUS

(9) Farbenfabriken Bayer Ag; DE 890958 C CAPLUS

(10) Glen, R; WO 0027394 A 2000 CAPLUS

(12) Hoechst Marion Roussel de Gmbh; EP 0908456 A

1999

CAPLUS

(13) Hoechst Marion Roussel de Gmbh; DE 19756388 A

1999 CAPLUS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L33 ANSWER 2 OF 14 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

2000:725451 CAPLUS

DOCUMENT NUMBER:

133:286497

TITLE:

Immunomodulatory compositions and methods of use

thereof

INVENTOR(S):

Onderdonk, Andrew B.; Tzianabos, Arthur O.; Miller,

Robert J.; Calias, Pericles

PATENT ASSIGNEE(S):

Genzyme Corporations, USA

SOURCE:

PCT Int. Appl., 62 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.			KI	ND	DATE			APPLICATION NO.				э.	DATE						
										_	- -								
W	С	2000	0594	90	A.	2	2000	1012		W	200	00-ช	s908°	7	2000	0406			
WO 2000059490			A3		20010215														
		W:	ΑE,	AG,	AL,	AM,	AT,	AU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CR,	
			CU,	CZ,	DE,	DK,	DM,	DZ,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	GM,	HR,	HU,	

ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU,

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LV, MA, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE,
             SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZW,
             AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE,
             DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF,
             CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                                          US 1999-128177
                                                           P 19990406
PRIORITY APPLN. INFO.:
                          MARPAT 133:286497
OTHER SOURCE(S):
     The invention relates to immunomodulatory compns. and related methods.
     The immunomodulatory compns. are useful for the prevention of sepsis and
     the treatment and prevention of diseases assocd. with inflammation and/or
     NOS. CM-cellulose/N-ethyl-N'-(3-dimethylaminopropyl)urea formulations
are
     described.
IT
     32897-26-0 121007-41-8
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (immunomodulatory compns.)
RN
     32897-26-0 CAPLUS
     Urea, N-[3-(dimethylamino)propyl]-N'-ethyl- (9CI) (CA INDEX NAME)
CN
      0
EtNH-C-NH-(CH<sub>2</sub>)<sub>3</sub>-NMe<sub>2</sub>
RN
     121007-41-8 CAPLUS
     1-Propanaminium, 3-[[(ethylamino)carbonyl]amino]-N,N,N-trimethyl-, iodide
CN
     (9CI) (CA INDEX NAME)
EtNH-C-NH-(CH<sub>2</sub>)<sub>3</sub>-N+Me<sub>3</sub>
          I -
L33 ANSWER 3 OF 14 CAPLUS COPYRIGHT 2001 ACS
                          2000:368337 CAPLUS
ACCESSION NUMBER:
                          133:4656
DOCUMENT NUMBER:
                          Preparation of heteroarylpyrazoles as p38 kinase
TITLE:
                          inhibitors
INVENTOR(S):
                          Anantanarayan, Ashok; Clare, Michael; Collins, Paul
                          W.; Crich, Joyce Z.; Devraj, Rajesh; Flynn, Daniel
L.;
                          Geng, Lifeng; Graneto, Matthew J.; Hanau, Cathleen
E.;
                          Hanson, Gunnar J.; Hartmann, Susan J.; Hepperle,
                          Michael; Huang, He; Khanna, Ish K.; Koszyk, Francis
```

J.; Liao, Shuyuan; Metz, Suzanne; Partis, Richard A.; Perry, Thao D.; Rao, Shashidhar N.; Selness, Shaun Raj; South, Michael S.; Stealey, Michael A.; et al.

1

PATENT ASSIGNEE(S):

G.D. Searle & Co., USA

SOURCE:

PCT Int. Appl., 1210 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE _____ _____ -----WO 2000031063 20000602 WO 1999-US26007 19991117 **A**1 W: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU, CZ, DE, DK, DM, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG PRIORITY APPLN. INFO.: US 1998-196623 A 19981120 MARPAT 133:4656

OTHER SOURCE(S):

Title compds. [I; R1 = H, OH, NH2, (cyclo)alk(en)yl, acyl, aryl, etc.; R2 = H, halo, alkyl, alkoxy, (un) substituted piperidinyl, etc.; R3 = pyridyl,

pyrimidinyl, quinolyl, etc.; R4 = H, alkyl, heterocyclyl, aryl, etc.] were

prepd. by reaction of ketones with hydrazines. Thus, R3CH2COMe (R3 = 4-pyridinyl) was condensed with 3,4-F(MeO)C6H3CHO and the product cyclocondensed with TsNHNH2 to give title compd. II. Data for biol. activity of I were given.

IT 216523-08-9P 216523-09-0P

> RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of heteroarylpyrazole p38 kinase inhibitors by cyclocondensation of hydrazines with ketones)

RN 216523-08-9 CAPLUS

CN

N-[2-(dimethylamino)ethyl]-N'-[3-[5-(4-fluorophenyl)-4-(4-pyridinyl)-4-(4-py1H-pyrazol-3-yl]propyl]- (9CI) (CA INDEX NAME)

$$Me_2N-CH_2-CH_2-NH-C-NH-(CH_2)_3$$

CN Urea, N-[3-(dimethylamino)propyl]-N'-[3-[5-(4-fluorophenyl)-4-(4-pyridinyl)-1H-pyrazol-3-yl]propyl]- (9CI) (CA INDEX NAME)

REFERENCE COUNT:

REFERENCE(S):

10

(1) Anantanarayan, A; WO 9852937 A 1998 CAPLUS

(2) Anantanarayan, A; WO 9852940 A 1998 CAPLUS

(3) Fujisawa Pharmaceutical Co; EP 0531901 A 1993 CAPLUS

(4) Lilly Co Eli; EP 0846687 A 1998 CAPLUS

(5) Oku Teruo; WO 9419350 A 1994 CAPLUS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L33 ANSWER 4 OF 14 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

2000:307141 CAPLUS

DOCUMENT NUMBER:

132:331676

TITLE:

Fluorescence immunoassays using analyte

(analog)-conjugated porphyrin-silicon complex fluorescent dyes free of aggregation and serum

binding

INVENTOR(S):

Devlin, Robert Francis; Dandliker, Walter Beach;

Arrhenius, Peter Olaf Gustaf

PATENT ASSIGNEE(S):

Hyperion, Inc., USA

SOURCE:

U.S., 58 pp., Cont.-in-part of U.S. 5,880,287.

CODEN: USXXAM

DOCUMENT TYPE:

Patent

LANGUAGE:

English

9

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 6060598	A	20000509	US 1997-874820	19970613
US 5403928	Α	19950404	US 1991-701449	19910515
US 5641878	Α	19970624	US 1994-333603	19941102
US 5677199	A	19971014	US 1994-346098	19941129
US 5880287	А	19990309	US 1995-476544	19950606
PRIORITY APPLN.	INFO.:		US 1990-523601 B2	19900515
			US 1990-524212 B2	19900515
			US 1991-701449 A3	19910515
			US 1991-701465 B1	19910515
			US 1994-333603 A2	19941102
			US 1994-346098 A2	19941129
			US 1995-476544 A2	19950606

AB Fluorescence immunoassay methods are provided which use fluorescent dyes which are free of aggregation and serum binding. Such immunoassay methods

are thus, particularly useful for the assay of biol. fluids, such as serum, plasma, whole blood and urine. The compds. of the invention, whose

prepn. is described, include silicon complexes with porphyrin derivs. which are linked to an analyte or analog thereof, e.g. a caged dicarboxy silicon phthalocyanine digoxin probe.

IT 267422-47-9P

RL: ARG (Analytical reagent use); BUU (Biological use, unclassified); PRP (Properties); SPN (Synthetic preparation); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); PREP (Preparation); USES (Uses)

(fluorescence immunoassays using analyte (analog)-conjugated porphyrin-silicon complex fluorescent dyes free of aggregation and serum binding)

RN 267422-47-9 CAPLUS

CN Poly(oxy-1,2-ethanediy1), .alpha.-hydro-.omega.-methoxy-, ester with hydrogen (OC-6-13)-[3-[[[(5.beta.,12.beta.,14.beta.)-21,23-epoxy-12,14-dihydroxy-23-oxo-24-norchol-20(22)-en-3-yl]amino]carbonyl]-29H,31H-

phthalocyanine-2-carboxylato(3-)-.kappa.N29,.kappa.N30,.kappa.N31,.kappa.N

32]bis[[2-[[[[3-[(hydroxy-.kappa.0)dimethylsilyl]propyl]amino]carbonyl]aminoll]ethyl]carbamato]silicate(1-) (2:1) (9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 1-B

PAGE 1-C

PAGE 2-A

● H+

PAGE 2-B

IT 267422-48-0P 267422-49-1P 267422-50-4P 267422-51-5P 267422-52-6P 267422-53-7P 267422-54-8P

RL: ARG (Analytical reagent use); BUU (Biological use, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); PREP (Preparation); USES (Uses) (fluorescence immunoassays using analyte (analog)-conjugated porphyrin-silicon complex fluorescent dyes free of aggregation and serum binding)

RN 267422-48-0 CAPLUS

CN Poly(oxy-1,2-ethanediyl), .alpha.-hydro-.omega.-methoxy-, ester with hydrogen

(OC-6-13)-[3-[[[(5.beta.,14.beta.)-21,23-epoxy-14-hydroxy-23-oxo-24-norchol-20(22)-en-3-yl]aminol]carbonyl]-29H,31H-phthalocyanine-2-carboxylato(3-)-.kappa.N29,.kappa.N30,.kappa.N31,.kappa.N32]bis[[2-[[[[3-

[(hydroxy-.kappa.O)dimethylsilyl]propyl]amino]carbonyl]aminol]ethyl]carbam ato]silicate(1-) (2:1) (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} \text{MeO} & \begin{array}{c|c} \text{CH}_2 - \text{CH}_2 - \text{O} & \begin{array}{c} \text{O} & \text{O} \\ \text{\parallel} & \text{C} - \text{NH} - \text{CH}_2 - \text{CH}_2 - \text{NH} - \text{C} - \text{NH} - \text{C}$$

PAGE 1-C

PAGE 2-A

● H+

__/

PAGE 2-B

RN 267422-49-1 CAPLUS
CN Poly(oxy-1,2-ethanediyl), .alpha.-hydro-.omega.-methoxy-, 1,1'-diester with trihydrogen
(OC-6-13)-[3-[[[2-[[4-(2,3,6,7,8,9-hexahydro-1,3-dimethyl-2,6-dioxo-1H-purin-8-yl)-1-oxobutyl]amino]ethyl]amino]carbonyl]-29H,31H-

phthalocyanine-2-carboxylato(3-)-.kappa.N29,.kappa.N30,.kappa.N31,.kappa.N
32]bis[11-(hydroxy-.kappa.O)-11-methyl-6-oxo-2,5,7-triaza-11siladodecanoato(2-)]silicate(3-) (9CI) (CA INDEX NAME)

PAGE 1-B

PAGE 2-A

● H+

PAGE 2-B

RN 267422-50-4 CAPLUS

CN Poly(oxy-1,2-ethanediyl), .alpha.-hydro-.omega.-methoxy-, 1,1'-diester with trihydrogen (OC-6-13)-[3-[[[4-(5-ethylhexahydro-2,4,6-trioxo-5-

pyrimidinyl)phenyl]amino]carbonyl]-29H,31H-phthalocyanine-2-carboxylato(3-

-)-.kappa.N29,.kappa.N30,.kappa.N31,.kappa.N32]bis[11-(hydroxy-.kappa.O)-11-methyl-6-oxo-2,5,7-triaza-11-siladodecanoato(2-)]silicate(3-) (9CI) (CA INDEX NAME)
- *** STRUCTURE DIAGRAM IS NOT AVAILABLE ***
- RN 267422-51-5 CAPLUS
- CN Poly(oxy-1,2-ethanediyl), .alpha.-hydro-.omega.-methoxy-, 1,1'-diester with trihydrogen
- (OC-6-13)-[3-[[[2-[[[4-(4-hydroxy-3,5-diiodophenoxy)-3,5-diiodophenyl]acetyl]amino]ethyl]amino]carbonyl]-29H,31H-phthalocyanine-2-carboxylato(3-)-.kappa.N29,.kappa.N30,.kappa.N31,.kappa.N32]bis[11-

(hydroxy-.kappa.O)-11-methyl-6-oxo-2,5,7-triaza-11-siladodecanoato(2)]silicate(3-) (9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 1-B

PAGE 1-C

PAGE 2-A

● H+

PAGE 2-B

RN 267422-52-6 CAPLUS

CN Poly(oxy-1,2-ethanediyl), .alpha.-hydro-.omega.-methoxy-, 1,1'-diester with trihydrogen

(OC-6-13)-[3-[[[2-[[4-(acetylamino)benzoyl]amino]ethyl]et hylamino]carbonyl]-29H,31H-phthalocyanine-2-carboxylato(3-)-.kappa.N29,.kappa.N30,.kappa.N31,.kappa.N32]bis[11-(hydroxy-.kappa.O)-11methyl-6-oxo-2,5,7-triaza-11-siladodecanoato(2-)]silicate(3-) (9CI) (CA INDEX NAME)

*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***

RN 267422-53-7 CAPLUS

CN Poly(oxy-1,2-ethanediy1), .alpha.-hydro-.omega.-methoxy-, 1,1'-diester with trihydrogen (OC-6-13)-[3-[[[4-(5-ethylhexahydro-4,6-dioxo-5-

pyrimidinyl)phenyl]amino]carbonyl]-29H,31H-phthalocyanine-2-carboxylato(3-

```
)-.kappa.N29,.kappa.N30,.kappa.N31,.kappa.N32]bis[11-(hydroxy-.kappa.O)-11-
     methyl-6-oxo-2,5,7-triaza-11-siladodecanoato(2-)]silicate(3-) (9CI) (CA
     INDEX NAME)
*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***
     267422-54-8 CAPLUS
     Poly(oxy-1,2-ethanediyl), .alpha.-hydro-.omega.-methoxy-, ester with
CN
     dihydrogen (OC-6-13)-[3-[[(carboxydiphenylmethyl)amino]carbonyl]-29H,31H-
phthalocyanine-2-carboxylato(4-)-.kappa.N29,.kappa.N30,.kappa.N31,.kappa.N
32]bis[[2-[[[[3-[(hydroxy-.kappa.0)dimethylsilyl]propyl]amino]carbonyl]ami
     no]ethyl]carbamato]silicate(2-) (2:1) (9CI) (CA INDEX NAME)
*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***
REFERENCE COUNT:
                          48
REFERENCE(S):
                          (1) Anon; WO 9118006 1981 CAPLUS
                          (2) Anon; EP 0260098 1987 CAPLUS
                          (5) Anon; JP 63264674 1988 CAPLUS
                          (6) Anon; EP 0336879 1989 CAPLUS
                          (7) Anon; WO 9002747 1990 CAPLUS
                          ALL CITATIONS AVAILABLE IN THE RE FORMAT
L33 ANSWER 5 OF 14 CAPLUS COPYRIGHT 2001 ACS
ACCESSION NUMBER:
                          1999:795794 CAPLUS
DOCUMENT NUMBER:
                          132:35701
TITLE:
                          Preparation of imidazolyl derivatives as as agonists
                          or antagonists of somatostatin receptors
                          Thurieau, Christophe Alain; Poitout, Lydie Francine;
INVENTOR(S):
                          Galcera, Marie-Odile; Gordon, Thomas D.; Morgan,
                          Barry; Moinet, Christophe Philippe
PATENT ASSIGNEE(S):
                          Societe de Conseils de Recherches et d'Applications
                          Scientifiques, S.A., Fr.
SOURCE:
                          PCT Int. Appl., 342 pp.
                          CODEN: PIXXD2
DOCUMENT TYPE:
                          Patent
LANGUAGE:
                          English
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
    PATENT NO.
                      KIND
                            DATE
                                            APPLICATION NO.
                      A2
                             19991216
                                            WO 1999-US12760 19990608
         W: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ,
             DE, DK, EE, ES, FI, GB, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP,
             KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN,
             MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM,
             TR, TT, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD,
             RU, TJ, TM
         RW: GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG,
             CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
    AU 9944257
                       A1
                            19991230
                                            AU 1999-44257
                                                              19990608
    EP 1086086
                            20010328
                                            EP 1999-927323
                       A1
                                                              19990608
```

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, PT, IE,

FI

NO 2000006267 A 20010207 PRIORITY APPLN. INFO.:

NO 2000-6267 20001211 US 1998-89087 P 19980612 US 1998-96431 A1 19980612 WO 1999-US12760 W 19990608

OTHER SOURCE(S): MARPAT 132:35701

The title compds. [I; R1 = H, (CH2)mCO(CH2)mZ1, (CH2)mZ1, etc.; Z1 = (un)substituted benzo[b]thiophene, Ph, naphthyl, etc.; R2 = H, alkyl; R1 and R2 taken together with the nitrogen atoms to which they are attached form II-IV; R3 = (CH2)mE(CH2)mZ2; E = O, S, CO, etc.; Z2 = H, alkyl, NH2, etc.; R4 = H, (CH2)mA1; A1 = C(:Y)NX1X2; C(:Y)X2; C(:NH)X2, X2; Y = O, S; X1 = H, alkyl, etc.; X2 = alkyl, etc.; R5 = alkyl, (un)substituted aryl, etc.; R6 = H, alkyl; R7 = alkyl, (CH2)mZ4; Z4 = (un)substituted Ph, naphthyl, indolyl, etc.; m = 0-6] which are useful as agonists or antagonists of somatostatin receptors (no data), and for inhibiting the proliferation of Helicobacter pylori, were prepd. Thus, activating 2-furancarboxylic acid with carbonyldiimidazole followed by addn. of 2-{(1S)-1-amino-2-(indol-3-yl)ethyl}-4-phenyl-1H-imidazole afforded 94% the title compd. V. Compds. I are effective at 0.01-10.0 mg/kg/day.

IT 252305-00-3P 252311-37-8P 252311-82-3P 252314-08-2P 252314-32-2P

RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of imidazolyl derivs. as as agonists or antagonists of somatostatin receptors)

RN 252305-00-3 CAPLUS

CN Urea, N-[(1S)-5-amino-1-(4-phenyl-1H-imidazol-2-yl)pentyl]-N'-[3-(dimethylamino)propyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 252311-37-8 CAPLUS

CN Urea, N-[3-(dimethylamino)propyl]-N'-[(1R)-2-(1H-indol-3-yl)-1-(4-phenyl-1H-imidazol-2-yl)ethyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 252311-82-3 CAPLUS

CN Urea,

N-[3-(dimethylamino)propyl]-N'-[1-[4-(1,1-dimethylethyl)-1H-imidazol-2-yl]-2-(1H-indol-3-yl)ethyl]-(9CI) (CA INDEX NAME)

$$\begin{array}{c|c} H & O \\ \parallel & NH-C-NH-(CH_2)_3-NMe_2 \\ \hline \\ CH_2-CH-N-Bu-t \\ N \end{array}$$

RN 252314-08-2 CAPLUS

CN Urea,

 $\begin{array}{lll} N-[2-(diethylamino)ethyl]-N'-[(1R)-2-(1H-indol-3-yl)-1-(4-phenyl-1H-imidazol-2-yl)ethyl]-& (9CI) & (CA INDEX NAME) \end{array}$

Absolute stereochemistry.

RN 252314-32-2 CAPLUS

CN Urea,

N-[2-(diethylamino)ethyl]-N'-[(1S)-2-(1H-indol-3-yl)-1-(4-phenyl-1H-imidazol-2-yl)ethyl]-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

```
L33 ANSWER 6 OF 14 CAPLUS COPYRIGHT 2001 ACS
```

ACCESSION NUMBER: 1999:783937 CAPLUS

DOCUMENT NUMBER: 132:22973

TITLE: Preparation of pyrrolo[2,3-d]pyrimidines as adenosine

receptor antagonists

INVENTOR(S): Castelhano, Arlindo L.; McKibben, Bryan; Witter,

David

Me)(II;

PATENT ASSIGNEE(S): Cadus Pharmaceutical Corp., USA

SOURCE: PCT Int. Appl., 169 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

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PATENT NO.
                     KIND DATE
                                          APPLICATION NO.
                                                            DATE
                            19991209
                                          WO 1999-US12135 19990601
    WO 9962518
                      A1
        W: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ,
            DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS,
             JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK,
            MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ,
            TM, TR, TT, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ,
            MD, RU, TJ, TM
        RW: GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW, AT, BE, CH, CY, DE, DK,
            ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG,
            CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
    AU 9942265
                      A1 19991220
                                          AU 1999-42265
                                                            19990601
                                          BR 1999-11612
    BR 9911612
                           20010206
                      Α
                                                            19990601
    EP 1082120
                                          EP 1999-926107
                           20010314
                      A1
                                                            19990601
            AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
            IE, SI, LT, LV, FI, RO
    NO 2000006090
                           20010131
                                          NO 2000-6090
                     Α
                                                            20001130
PRIORITY APPLN. INFO.:
                                        US 1998-87702
                                                         Р
                                                            19980602
                                        US 1999-123216
                                                         Ρ
                                                            19990308
                                        US 1999-126527
                                                         Ρ
                                                            19990326
                                        WO 1999-US12135 W
                                                            19990601
OTHER SOURCE(S):
                        MARPAT 132:22973
    Title compds. [I; R = NR1R2; R1-R4 = H, alkyl, aryl, etc.; NR1R2 =
    heterocyclyl; R5,R6 = H, halo, alkyl, aryl, etc.; R4R5,R5R6 = atoms to
    complete a ring) were prepd. Thus, 2- amino-3-cyano-4,5-dimethyl-1-(1-
    phenylethyl)pyrrole was N-benzoylated and the product cyclized to give,
    after deprotection and chlorination, I (R3 = Ph, R4 = H, R5 = R6 =
```

R = Cl) which was aminated by trans-4-hydroxycyclohexylamine to give II

(R

= trans-4-hydroxycyclohexylamino). Data for biol. activity of I were given.

IT 251946-33-5P 251946-34-6P

RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of pyrrolo[2,3-d]pyrimidines as adenosine receptor
antagonists)

RN 251946-33-5 CAPLUS

CN Urea, N-[2-[(5,6-dimethyl-2-phenyl-1H-pyrrolo[2,3-d]pyrimidin-4-yl)amino]ethyl]-N'-methyl- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & & H & & Me \\ & & & & N & & H & \\ & & & & N & & Me \\ \\ & & & & & MeNH-C-NH-CH_2-CH_2-NH & & & \\ \end{array}$$

RN 251946-34-6 CAPLUS

CN Urea, N-[2-[(5,6-dimethyl-2-phenyl-1H-pyrrolo[2,3-d]pyrimidin-4-yl)amino]ethyl]-N'-ethyl- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & H & Me \\ & & & N & Me \\ & & & \\ & & \\ & & & \\ & & & \\ & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & &$$

REFERENCE COUNT:

20

REFERENCE(S):

- (1) Chen Yuhpyng, L; WO 9413676 A 1994 CAPLUS
- (2) Ciba Geigy AG; EP 0682027 A 1995 CAPLUS
- (3) Hitchings, G; US 3037980 A 1962 CAPLUS
- (4) Hoechst India Ltd; IN 157280 A 1986 CAPLUS
- (5) Iwamura, H; J Med Chem 1983, V26, P838 CAPLUS
- ALL CITATIONS AVAILABLE IN THE RE FORMAT

L33 ANSWER 7 OF 14 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

1999:763780 CAPLUS

DOCUMENT NUMBER:

132:10496

TITLE:

Method for preparing thin liquid samples for

microscopic analysis

INVENTOR(S):

Berndt, Klaus W.

PATENT ASSIGNEE(S):

Becton, Dickinson and Company, USA

SOURCE:

Eur. Pat. Appl., 14 pp. CODEN: EPXXDW

DOCUMENT TYPE:

Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

EP 961109 A2 19991201 EP 1999-108936 19990505

EP 961109 A3 20000719

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO

JP 2000002839 A2 20000107 JP 1999-147168 19990526 PRIORITY APPLN. INFO.: US 1998-85851 A 19980527

AB A method for producing thin samples of liqs. for microscopic anal. involves depositing a drop of the liq. sample onto the upper surface of a microscope slide near the center of the slide, positioning a flexible cover glass onto spacers on the slide, applying a downward force to the upper surface of the cover glass so that the lower surface of the cover glass touches the sample, suspending the application of force, and obtaining a thin liq. sample. A liq. blood sample prepd. this way had a central area A contg. plasma but no red blood cells. This region A was surrounded by a wide ring B contg. huge nos. of isolated red blood cells in a well-defined monolayer arrangement. Ring B was surrounded by an

even

wider belt that contained red blood cells in Rouleaux formation where the length of the Rouleaux blocks increased with increasing distance from the center. This kind of blood sample prepn. does not result in morphol. changes as obsd. in the wedge slide method or during drying of blood

films in the open air.

IT 154088-80-9, LaJolla Blue

RL: ARG (Analytical reagent use); DEV (Device component use); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); USES (Uses)

(deposited on microscope slide; method for prepg. thin liq. samples

for

microscopic anal.)

RN 154088-80-9 CAPLUS

CN Poly(oxy-1,2-ethanediyl), .alpha.-hydro-.omega.-methoxy-, ether with dihydrogen (OC-6-12)-bis(2-hydroxyethyl 11-hydroxy-11-methyl-6-oxo-2,5,7-

triaza-11-siladodecanoato-011)[29H,31H-phthalocyanine-2,3-carboxylato(4-)-N29,N30,N31,N32]silicate(2-) (2:1) (9CI) (CA INDEX NAME)

*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***

L33 ANSWER 8 OF 14 CAPLUS COPYRIGHT 2001 ACS ACCESSION NUMBER: 1998:789144 CAPLUS

DOCUMENT NUMBER:

130:38377

TITLE:

Preparation of heteroarylpyrazoles as p38 kinase

inhibitors

INVENTOR(S):

Anantanarayan, Ashok; Clare, Michael; Collins, Paul W.; Crich, Joyce Zuowu; Devraj, Rajesh; Flynn, Daniel L.; Geng, Lifeng; Hanson, Gunnar J.; Koszyk, Francis

J.; Liao, Shuyuan; Partis, Richard A.; Rao,

Shashidhar

N.; Selness, Shaun Raj; South, Michael S.; Stealey,

```
Michael A.; Weier, Richard M.; Xu, Xiangdong; et al.
PATENT ASSIGNEE(S):
                         G.D. Searle and Co., USA; et al.
SOURCE:
                         PCT Int. Appl., 828 pp.
                         CODEN: PIXXD2
DOCUMENT TYPE:
                         Patent
LANGUAGE:
                         English
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:
     PATENT NO.
                     KIND DATE
                                        APPLICATION NO. DATE
     WO 9852940 A1 19981126 WO 1998-US10436 19980522
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             KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX,
             NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT,
             UA, UG, US, UZ, VN, YU, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES,
             FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI,
             CM, GA, GN, ML, MR, NE, SN, TD, TG
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                                        AU 1998-75883
                                                           19980522
                          19990524
     ZA 9804358
                      Α
                                          ZA 1998-4358
                                                           19980522
                     A1 20000517
     EP 1000055
                                         EP 1998-923642
                                                           19980522
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
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                                          BR 1998-9147
                                                           19980522
     NO 9905695
                      Α
                           20000121
                                          NO 1999-5695
                                                           19991119
PRIORITY APPLN. INFO.:
                                       US 1997-47570
                                                        Ρ
                                                           19970522
                                       WO 1998-US10436 W
                                                          19980522
OTHER SOURCE(S):
                       MARPAT 130:38377
    Title compds. [I; R1 = H, NH2, (cyclo)alk(en)yl, acyl, aryl, etc.; R2 =
Η,
     halo, alkyl, alkoxy, etc.; R3 = pyridyl, pyrimidinyl, quinolyl, etc.; R4
     H, alkyl, heterocyclyl, aryl, etc.] were prepd. Thus, R3CH2COMe (R3 =
     4-pyridinyl) was condensed with 3,4-F(MeO)C6H3CHO and the product
     cyclocondensed with TsNHNH2 to give title compd. II. Data for biol.
     activity of I were given.
     216523-08-9P 216523-09-0P
    RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic
     preparation); THU (Therapeutic use); BIOL (Biological study);
     PREP (Preparation); USES (Uses)
        (prepn. of heteroarylpyrazoles as p38 kinase inhibitors)
     216523-08-9 CAPLUS
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L6
L7
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L8
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L9
              4 S ETHYL(L) DIMETHYLAMINOPROPYL(L) UREA
L10
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L16
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     FILE 'BEILSTEIN' ENTERED AT 10:57:15 ON 01 JUN 2001
L17
              1 S L11
     FILE 'REGISTRY' ENTERED AT 10:57:49 ON 01 JUN 2001
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L20
L21
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L22
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L24
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    FILE 'USPATFULL' ENTERED AT 11:03:35 ON 01 JUN 2001
L25
             7 S L22
    FILE 'REGISTRY' ENTERED AT 11:05:41 ON 01 JUN 2001
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L26
L27
                STRUCTURE UPLOADED
L28
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                STRUCTURE UPLOADED
L29
L30
           1523 S L29 FULL SUB=L28
L31
           381 S L30 AND 1/0
L32
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FULL ESTIMATED COST
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CA SUBSCRIBER PRICE
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STRUCTURE FILE UPDATES: 30 MAY 2001 HIGHEST RN 339046-06-9 DICTIONARY FILE UPDATES: 30 MAY 2001 HIGHEST RN 339046-06-9

TSCA INFORMATION NOW CURRENT THROUGH January 11, 2001

Please note that search-term pricing does apply when conducting SmartSELECT searches.

Structure search limits have been increased. See HELP SLIMIT for details.

=> s 132 and 1/nr 2662225 1/NR L34 49 L32 AND 1/NR

=> d scan

L34 49 ANSWERS REGISTRY COPYRIGHT 2001 ACS
IN Urea, N-[2-(dimethylamino)ethyl]-N'-(3-phenylpropyl)- (9CI)
MF C14 H23 N3 O

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):1

L34 49 ANSWERS REGISTRY COPYRIGHT 2001 ACS
IN Urea, N-[(diethylamino)methyl]-N'-[(2-methyl-4-pyridinyl)methyl]- (9CI)
MF C13 H22 N4 O

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

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ENTRY SESSION
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DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)
SINCE FILE TOTAL
ENTRY SESSION
SESSION

CA SUBSCRIBER PRICE

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FILE COVERS 1947 - 1 Jun 2001 VOL 134 ISS 23 FILE LAST UPDATED: 30 May 2001 (20010530/ED)

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This file supports REG1stRY for direct browsing and searching of all substance data from the REGISTRY file. Enter HELP FIRST for more information.

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     FILE 'CAPLUS' ENTERED AT 11:01:43 ON 01 JUN 2001
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     FILE 'USPATFULL' ENTERED AT 11:03:35 ON 01 JUN 2001
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              7 S L22
     FILE 'REGISTRY' ENTERED AT 11:05:41 ON 01 JUN 2001
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L27
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L28
           3145 S L27 FULL SUB=L26
L29
                STRUCTURE UPLOADED
L30
           1523 S L29 FULL SUB=L28
L31
            381 S L30 AND 1/0
L32
            331 S L31 NOT S/ELS
     FILE 'CAPLUS' ENTERED AT 11:14:57 ON 01 JUN 2001
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19362000 PY>=199

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09/350,193
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L37 ANSWER 1 OF 26 CAPLUS COPYRIGHT 2001 ACS
ACCESSION NUMBER:
                          1998:220203 CAPLUS
DOCUMENT NUMBER:
                          129:4517
TITLE:
                          Solid phase organic synthesis of polyamine
derivatives
                          and initial biological evaluation of their
antitumoral
                          activity
AUTHOR(S):
                          Tomasi, Sophie; Le Roch, Myriam; Renault, Jacques;
                          Corbel, Jean-Charles; Uriac, Philippe; Carboni,
                          Bertrand; Moncoq, Damien; Martin, Benedicte; Delcros,
                          Jean-Guy
CORPORATE SOURCE:
                          Pharmacochimie de Molecules de Synthese et de
Produits
                         Naturels, Fac. de Pharmacie, Rennes, 35043, Fr.
SOURCE:
                         Bioorg. Med. Chem. Lett. (1998), 8(6), 635-640
                         CODEN: BMCLE8; ISSN: 0960-894X
PUBLISHER:
                         Elsevier Science Ltd.
DOCUMENT TYPE:
                         Journal
LANGUAGE:
                         English
     A series of N1-monosubstituted putrescine and spermine derivs. was
     synthesized using a solid phase methodol. Their cytotoxicity, calmodulin
     antagonism and polyamine uptake inhibition, pharmacol. properties shared
     by some antitumoral agents was evaluated.
     207501-42-6P
     RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic
     preparation); BIOL (Biological study); PREP (Preparation)
        (solid phase org. synthesis of polyamine derivs. and initial biol.
        evaluation of antitumoral activity)
RN
     207501-42-6 CAPLUS
CN
     Urea,
N-[3-[[4-[(3-aminopropyl)amino]butyl]amino]propyl]-N'-(phenylmethyl)-
     , trifluoroacetate (9CI) (CA INDEX NAME)
     CM
          1
     CRN 207501-41-5
     CMF C18 H33 N5 O
Ph-CH_2-NH-C-NH-(CH_2)_3-NH-(CH_2)_4-NH-(CH_2)_3-NH_2
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CRN 76-05-1 CMF C2 H F3 O2

L37 ANSWER 2 OF 26 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 1997:366218 CAPLUS

DOCUMENT NUMBER: 127:95010

TITLE: Selective synthesis of polyamine derivatives.

Efficient derivatization of the secondary amino group

of N-monosubstituted 1,3-diamines

AUTHOR(S): Jentgens, Christian; Hofmann, Richard; Guggisberg,

Armin; Bienz, Stefan; Hesse, Manfred

CORPORATE SOURCE: Organisch-Chemisches Inst., Universitat Zurich,

Zurich, CH-8057, Switz.

SOURCE: Helv. Chim. Acta (1997), 80(3), 966-978

CODEN: HCACAV; ISSN: 0018-019X Verlag Helvetica Chimica Acta

DOCUMENT TYPE: Journal

LANGUAGE: English

OTHER SOURCE(S): CASREACT 127:95010

AB N-monosubstituted 1,3-diamines were selectively functionalized at the secondary N atom via 2-phenyl-substituted hexahydropyrimidine intermediates. Reaction of the diamines with PhCHO, followed by

treatment

PUBLISHER:

with an electrophile and hydrolysis, provided the desired products with excellent selectivity and in high yields. N4,N9-bis[3-phenylprop-2-enoyl]spermine (I), which was further converted to

N1, N12-bis[3-phenylprop-

2-enoyl]spermine by a transamidation reaction, was prepd. by this way in 82% yield from spermine. Compd. I was alternatively synthesized in 83% yield, equally from spermine, by a sequence involving intermediary protection of the terminal amino groups.

IT 191990-75-7P

RL: SPN (Synthetic preparation); PREP (Preparation)

(prepn. of polyamines by selective derivatization of secondary amino group of monosubstituted diamines)

RN 191990-75-7 CAPLUS

CN Urea, N-[3-(methylamino)propyl]-N'-(phenylmethyl)- (9CI) (CA INDEX NAME)

O || MeNH- (CH₂)₃-NH-C-NH-CH₂-Ph

L37 ANSWER 3 OF 26 CAPLUS COPYRIGHT 2001 ACS ACCESSION NUMBER: 1996:476785 CAPLUS

DOCUMENT NUMBER:

125:142463

CODEN: EPXXDW

TITLE:

INVENTOR(S):

Carbodiimide derivatives for use in biotinylations

Takenishi, Soichiro; Suzuki, Osamu; Yokomizo,

Hirohiko; Ichihara, Tatsuo; Masuda, Gen; Shiohata,

Namiko; Komiya, Kazuko

PATENT ASSIGNEE(S):

Nisshinbo Industries, Inc., Japan Eur. Pat. Appl., 55 pp.

SOURCE:

Patent

DOCUMENT TYPE: LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE	
EP 718300	A1	19960626	EP 1995-309433	19951222	
R: DE, FR, JP 08176159	GB A2	19960709	JP 1994-335492	19941222	
US 5700935	Α	19971223	US 1995-577374	19951222	
US 5789588	Α	19980804	US 1997-931714	19970916	
PRIORITY APPLN. INFO	.:		JP 1994-335492	19941222	
			US 1995-577374	19951222	

OTHER SOURCE(S):

MARPAT 125:142463

Carbodiimides W1-X-N=C=N-Y-W2-Z [W1 = aliph., (un) substituted aryl, heteroaryl, tertiary amino, quaternary ammonium; -W2-Z = quaternary ammonium; X and Y = bond, alkylene; Z = biotin-contg. group] are useful

as

labeling reagents for introducing a biotin group into a nucleic acid or a protein. Thus, cyclohexyl isocyanate was treated with Me2NC6H4NH2-4 to give the urea which was converted to the carbodiimide and treated with 6-iodohexylbiotinamide to give the quaternized deriv. I.

179540-21-7P 179540-28-4P 179540-73-9P ΙT 179540-75-1P 179540-96-6P 179541-13-0P 179541-47-0P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn. of carbodiimide derivs. of biotin for use in biotinylations)

RN 179540-21-7 CAPLUS

CN Urea, N-[2-(dimethylamino)ethyl]-N'-(3-pyridinylmethyl)- (9CI) (CA INDEX NAME)

$$Me_2N-CH_2-CH_2-NH-C-NH-CH_2$$

RN 179540-28-4 CAPLUS

CN Urea, N-[(diethylamino)methyl]-N'-[(2-methyl-4-pyridinyl)methyl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c}
0 & N \\
\parallel & \parallel \\
\text{Et}_2\text{N}-\text{CH}_2-\text{NH}-\text{C}-\text{NH}-\text{CH}_2
\end{array}$$
Me

RN 179540-73-9 CAPLUS

CN Urea, N-[2-(diethylamino)ethyl]-N'-(2-pyridinylmethyl)- (9CI) (CA INDEX NAME)

RN 179540-75-1 CAPLUS

CN Urea, N-[3-(diethylamino)propyl]-N'-[(3-methyl-4-pyridinyl)methyl]- (9CI) (CA INDEX NAME)

RN 179540-96-6 CAPLUS

CN Urea, N-[(diethylamino)methyl]-N'-[2-(1-piperidinyl)ethyl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & \circ \\ & \parallel \\ & \land \\ & \land$$

RN 179541-13-0 CAPLUS

CN Urea, N-[2-(diethylamino)ethyl]-N'-[(1-ethyl-2-pyrrolidinyl)methyl]-(9CI)

(CA INDEX NAME)

RN 179541-47-0 CAPLUS

CN Urea, N-[2-(dimethylamino)ethyl]-N'-[2-(1-piperidinyl)ethyl]- (9CI) (CA INDEX NAME)

$$CH_2-CH_2-NH-C-NH-CH_2-CH_2-NMe_2$$

L37 ANSWER 4 OF 26 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 1995:451865 CAPLUS

DOCUMENT NUMBER: 122:214910

TITLE: Polyaniline derivatives and their manufacture

INVENTOR(S):
Oka, Osamu

PATENT ASSIGNEE(S): Tomoegawa Paper Co Ltd, Japan SOURCE: Jpn. Kokai Tokkyo Koho, 9 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

JP 06256510 A2 19940913 JP 1993-62428 19930301

AB Polyaniline derivs., which are gelatinizable and sol. in org. solvents, consist of polyaniline chains with no.-av. mol. wt. 2,000-500,000 and polyurea chains with no.-av. mol. wt. 180-100,000. Polyaniline chains

are

crosslinked by polyurea chains through amino groups of polyaniline. One such polymer was obtained by reaction polyaniline with an isocyanato-terminated polyurea made from 1,6-hexanediamine, urea, and phospene.

IT 161858-74-8P

RL: IMF (Industrial manufacture); PRP (Properties); PREP (Preparation) (polyaniline derivs. and their manuf.)

RN 161858-74-8 CAPLUS

CN Benzenamine, polymer with .alpha.-[9-[(4-chloro-1,4-dioxobutyl)amino]nonyl]-.omega.-[(4-chloro-1,4-

dioxobutyl)amino]poly(iminocarbonylimino-1,9-nonanediyl), graft (9CI)

(CA

INDEX NAME)

CM 1

CRN 161858-72-6

CMF (C10 H20 N2 O)n C17 H28 C12 N2 O4

CCI PMS

PAGE 1-B

CM 2

CRN 62-53-3 CMF C6 H7 N

L37 ANSWER 5 OF 26 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

1994:195651 CAPLUS

DOCUMENT NUMBER:

120:195651

TITLE:

Motor fuel detergent additives - asymmetrical ureas

of

hydrocarbyloxypolyether amines and tertiary

aminoalkyl

primary amines Herbstman, Sheldon PATENT ASSIGNEE(S): Texaco Inc., USA U.S., 8 pp.

SOURCE:

CODEN: USXXAM

DOCUMENT TYPE:

INVENTOR(S):

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. -----A 19940215 US 1992-910912 19920709

The present invention provides a novel class of compds., useful as AΒ gasoline detergent additives, comprising asym. ureas of either a hydrocarbyloxypolyether amine alone, or a hydrocarbyloxypolyether amine and a tertiary aminoalkyl primary amine. The present invention also provides a motor fuel compn. contg. the novel asym. ureas and further provides a method of synthesizing the asym. ureas of the present invention.

IT 153986-92-6P

RL: PREP (Preparation)

(prepn. of, gasoline detergent additive)

RN 153986-92-6 CAPLUS

CN Poly[oxy(methyl-1,2-ethanediyl)], .alpha.-[2-[[[[3-

(dimethylamino)propyl]amino]carbonyl]amino]-1-methylethyl]-.omega.-(4-

nonylphenoxy) - (9CI) (CA INDEX NAME)

PAGE 1-A

$$Me_{2}N-(CH_{2})_{3}-NH-C-NH-CH_{2}-CH-CH_{2}-O-(C_{3}H_{6})$$

PAGE 1-B

 \sim (CH₂)₈-Me

L37 ANSWER 6 OF 26 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

1993:168949 CAPLUS

DOCUMENT NUMBER:

118:168949

TITLE:

The preparation of N-alkyl-2-(1H)-pyridones by the

reaction of amines with a derivative of

3-(2-pyridyl)propane-1,2-diol

AUTHOR(S):

Block, Michael H.

CORPORATE SOURCE:

ICI Pharm., Mereside, Macclesfield/Cheshire, SK10

4TG,

UK

SOURCE:

Tetrahedron Lett. (1992), 33(52), 8149-50

CODEN: TELEAY; ISSN: 0040-4039

DOCUMENT TYPE:

Journal

LANGUAGE:

English

OTHER SOURCE(S):

CASREACT 118:168949

The redn. of the 2-pyridyl orthoester deriv. I with DIBAL gives exclusively the secondary alc. II (i.e. a 3-(2-pyridyl)propane-1,2-diol deriv.) in excellent yield. Mesylation of II followed by reaction with amines gives unusual N-alkyl-2(1H)-pyridones such as III. The prepn. of 1-amino-3-(2-pyridyloxy)-2-propanol derivs. by this method failed; the latter compds. are potential drugs for the treatment of congestive heart failure.

IT 71676-11-4

RL: RCT (Reactant)

(amination with, of (methoxymethoxy) (pyridyloy) propanol)

RN 71676-11-4 CAPLUS CN Urea, N-(2-aminoethyl)-N'-(phenylmethyl)- (9CI) (CA INDEX NAME)

О || Ph-CH₂-NH-C-NH-CH₂-CH₂-NH₂

L37 ANSWER 7 OF 26 CAPLUS COPYRIGHT 2001 ACS ACCESSION NUMBER: 1992:107071 CAPLUS

DOCUMENT NUMBER: 116:107071

TITLE: Copolymers with inherent antimicrobial activity

INVENTOR(S): Olstein, Alan D.

PATENT ASSIGNEE(S): Fuller, H. B., Licensing and Financing, Inc., USA

SOURCE: PCT Int. Appl., 33 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

W: CA, JP

RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LU, NL, SE

PRIORITY APPLN. INFO.: US 1990-479840 19900214

AB The title polymers contain .gtoreq.1 mol% unsatd. quaternary ammonium compds. and comonomers. Polymg.

m-CH2:C(Me)CH2C6H4CMe2NHCO2CH2CH2N+(C10H2

1) Me I-, Me methacrylate, Bu acrylate, and methacrylic acid in H2O at 65. degree. gave a 2.5:23.4:23.4:0.7 copolymer which was degraded by molds in 1 mo but resisted bacteria and yeasts.

IT **1393**62-81-5P

RL: RCT (Reactant); PREP (Preparation)
 (prepn. and quaternization of)

RN 139362-81-5 CAPLUS

CN Urea, N-[3-(dimethylamino)propyl]-N'-[1-methyl-1-[3-(1-methylethenyl)phenyl]ethyl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c}
 & \bigcirc \\
 & \parallel \\$$

L37 ANSWER 8 OF 26 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 1992:61651 CAPLUS

DOCUMENT NUMBER: 116:61651

TITLE: Copolymerizable imidazolidinones and oxazolidinones

INVENTOR(S):

Murdock, Thomas O.

PATENT ASSIGNEE(S):

Fuller, H. B., Licensing and Financing, Inc., USA

SOURCE:

PCT Int. Appl., 57 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	. DATE				
WO 9112243	A2	19910822	WO 1991-US939	19910212				
WO 9112243 W: CA, JP	A3	19911003						
RW: AT, BE		, DK, ES,	FR, GB, GR, IT, LU, NL	, SE				
CA 2074098	AA	19910815	CA 1991-2074098	19910212				
JP 05503941	Т2	19930624	JP 1991-504715	19910212				
EP 594596	A1	19940504	EP 1991-904976	19910212				
R: AT, BE	, CH, DE	, DK, ES,	FR, GB, GR, IT, LI, LU	, NL, SE				
PRIORITY APPLN. INF			US 1990-479718	19900214				
			WO 1991-US939	19910212				

MARPAT 116:61651

The title monomers, useful in coatings, caulks, sealing compns., adhesives, and as adhesion promoters, are prepd. Thus, I wain EtOAc at 25-30.degree. by dropwise addn. of II. An emulsion of copolymer from Bu acrylate 659, 2-ethylhexyl acrylate 90, methacrylic acid 20, methacrylonitrile 16, and I [prepd. from m-CH2:C(Me)C6H4C(Me2)NCO and 1-(2-aminoethyl)imidazolidin-2-one] 9.6% was used as a caulking compn. with better adhesion than without I.

IT137559-82-1P

RL: PREP (Preparation)

(coating emulsions, manuf. of, with good adhesion)

RN 137559-82-1 CAPLUS

CN2-Propenoic acid, 2-methyl-, polymer with butyl 2-propenoate, N-[2-(diethylamino)ethyl]-N'-[1-methyl-1-[3-(1-methyl-1)ethyl]-N'-[1-methyl-1-[3-(1-methyl-1)ethyl]-N'-[1-methyl-1-[3-(1-methyl-1)ethyl]-N'-[1-methyl-1-[3-(1-methyl-1)ethyl]-N'-[1-methyl-1-[3-(1-methyl-1)ethyl]-N'-[1-methyl-1-[3-(1-methyl-1)ethyl]-N'-[1-methyl-1-[3-(1-methyl-1)ethyl]-N'-[1-methyl-1-[3-(1-methyl-1)ethyl]-N'-[1-methyl-1-[3-(1-methyl-1)ethyl]-N'-[1-methyl-1-[3-(1-methyl-1)ethyl]-N'-[1-methyl-1-[3-(1-methyl-1)ethyl]-N'-[3-(1-memethylethenyl)phenyl]ethyl]urea and methyl 2-methyl-2-propenoate (9CI) (CA INDEX NAME)

CM 1

137559-81-0 CRN CMF C19 H31 N3 O

$$\begin{array}{c|c} & & & \\ & & & \\ \text{Me} - & & \\ & & \\ \text{CH}_2 & & \text{Me} \end{array}$$

09/350,193

CRN 141-32-2 CMF C7 H12 O2

$$\begin{array}{c} \text{O} \\ \parallel \\ \text{n-BuO-C-CH} \end{array} = \text{CH}_2$$

CM 3

CRN 80-62-6 CMF C5 H8 O2

$$\begin{array}{c|c} \text{H}_2\text{C} & \text{O} \\ \parallel & \parallel \\ \text{Me-C-C-OMe} \end{array}$$

CM 4

CRN 79-41-4 CMF C4 H6 O2

$$\begin{array}{c} \text{CH}_2 \\ || \\ \text{Me-C-CO}_2 \text{H} \end{array}$$

IT 137559-81-0P

RN 137559-81-0 CAPLUS

CN Urea, N-[2-(diethylamino)ethyl]-N'-[1-methyl-1-[3-(1-methylethenyl)phenyl]ethyl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & & \\ \text{Me} - & & \\ \text{C} & & & \\ & & & \\ \text{CH}_2 & & \text{Me} \end{array}$$

L37 ANSWER 9 OF 26 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

1991:2996 CAPLUS

DOCUMENT NUMBER:

114:2996

TITLE:

Direct-measuring assay dipsticks, their construction and use, and a dipstick-containing kit

INVENTOR(S):

Allen, Michael P.; Shibuya, Robert B.

PATENT ASSIGNEE(S):

Chemtrak, Inc., USA Eur. Pat. Appl., 16 pp.

SOURCE:

CODEN: EPXXDW

DOCUMENT TYPE:

Patent

LANGUAGE:

English

LANGUAGE:

די ידואויי

FAMILY ACC. NUM. COUNT:

Fudii

PATENT INFORMATION:

PA	TENT NO.		KIND	DATE	APPLICATION NO.	DATE
EΡ	342447		A2	19891123	EP 1989-108100	19890505
EΡ	34 2447		A3	19910731	1	
EP	342447		В1	19941214		
	R: AT,	BE,	CH, DE	, ES, FR, GB,	GR, IT, LI, LU, NL	, SE
US	4999287		Α	19910312	US 1988-195881	19880519
AU	89 33847		A1	19891123	AU 1989-33847	19890428
ΑU	6 26853		B2	19920813		
TD	02138961		A2	19900528	JP 1989-122996	19890518

PRIORITY APPLN. INFO.:

JP 1989-122996 19890518 US 1988-195881 19880519

AB The title dipsticks e.g. comprise, in the direction of fluid flow, (1) a 1st bibulous bridging strip extending from the immersion end to a sample pad site; (2) a sample pad; (3) a 2nd bibulous bridging strip extending from the sample pad site to a measurement region fluid receiving site, in which the 2nd strip is in liq. communication with the sample pad; (4) an extended bibulous measuring strip in communication with the 2nd bridging strip and impregnated with a 1st member of a signal-producing system, which upon reaction with a 2nd member of the signal-producing system produces a detectable signal defining a boundary on the measurement

strip;

and (5) a means of inhibiting fluid communication between the sample pad and the 1st and 2nd bridging strips prior to measurement and for permitting fluid communication during measurement. Means may also be induced for automatically metering sample vol. and for providing a sharply

delinessed color front. The method finds particular use where a limited amt. of substrate is provided for an enzyme on a sample pad. Thus, sample

strips were prepd. for detn. of 50-400 mg cholesterol equiv./dL in serum. The 3-methyl-2-benzothiazolinone hydrazone (mBTH) substrate was immobilized in the quantitation area at 0.25 or 0.50 mg/mL. Migration height of the color band was related to cholesterol concn., and the sensitivity of the assay was related to the amt. of immobilized mBTH.

IT 127931-32-2D, Whatman 3ET conjugates

RL: AN. 1 (Analytical study)

(in dipstick with discontinuous flow path for cholesterol detn.)

RN 127931-32-2 CAPLUS

CN Urea, N-(2-aminoethyl)-N'-[4-(methylphenylamino)butyl]- (9CI) (CA INDEX NAME)

```
L37 ANSWER 10 OF 26 CAPLUS COPYRIGHT 2001 ACS
ACCESSION NUMBER:
                        1989:440561 CAPLUS
DOCUMENT NUMBER:
                        111:40561
TITLE:
                        Preparation of diureidopolyaxylalkylene amine-blocked
                        isocyanate prepolymers for coatings
INVENTOR(S):
                        Speranza, George Phillip; Lin, Jiang Jen; Cuscurida,
                        Michael
PATENT ASSIGNEE(S):
                        Texaco Development Corp., USA
SOURCE:
                        Eur. Pat. Appl., 21 pp.
                        CODEN: EPXXDW
DOCUMENT TYPE:
                        Patent
LANGUAGE:
                        English
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
     PATENT NO. KIND DATE
                                        APPLICATION NO. DATE
     ______
    EP 301718 A2 A3
                                         ______
                                        EP 1988-306167 19880706
                           19890201
                          19891206
        R: DE, FR, GB
    US 4761465 A
                          19880802
                                         US 1987-78314
                                                         19870727
    US 4906774
                          19900306
                                         US 1987-78309
                     A
                                                         19870727
    CA 1328467
                     A1
                                         CA 1988-568200
                          19940412
                                                         19880531
    US 5010160
                     A
                          19910423
                                         US 1989-430686
                                                         19891030
PRIORITY APPLH. INFO.:
                                      US 1987-78309
                                                         19870727
                                      US 1987-78314
                                                         19870727
    The title polymers (mol. wt. 600-10,000) are prepd. from aliph.
    diisocy nates and polyoxyalkylene diamines in alc. solvents, or with
    excess diisocyanate and blocked with agents such as MEK oxime. Adding
0.1
    mol isophorane diisocyanate in 22.2 g iso-PrOH to 0.20 mol
    polyoxypropylene diamine (Jeffamine 400) over 1.5 h at 30.degree. gave a
    product with amine content 1.53 mequiv./g, while prepn. in (MeO) 2CO gave
а
    gel.
ΙT
    121467-12-9
    RL: US... (Uses)
       (coaring, impact-resistant)
RN
    121467-12-9 CAPLUS
    1,3-Probanediol, 2-ethyl-2-(hydroxymethyl)-, polymer with
    1,3-dii reganatomethylbenzene and .alpha.,.alpha.'-[1,15(1,16 or
    2,15) -c ...ethyl-4,13-dioxo-3,5,12,14-tetraazahexadecane-1,16-
    divl]bis[.omega.-(2-aminomethylethoxy)poly[oxy(methyl-1,2-ethanediyl)]]
    (9CI) (CA INDEX NAME)
    CM
         1
    CRN 11 382-85-2
    CMF ( CCI II
           H6 O)n (C3 H6 O)n C20 H44 N6 O4
             PMS
    CDES *
```

PAGE 1-A

$$H_2N-CH_2-CH_2-O$$
 $CH_2-CH_2-OH_2-CH_2-NH-C-NH-(CH_2)_6-$

4 (D1-Me)

PAGE 1-B

$$-NH-C-NH-CH2-CH2-CH2-CH2-CH2-CH2-NH2$$

CM 2

CRN 26471-62-5 CMF C9 H6 N2 O2 CCI IDS

CDES 8:ID

D1-Me

CM 3 •

CRN 77-99-6 CMF C6 H14 O3

L37 ANSWER 11 OF 26 CAPLUS COPYRIGHT 2001 ACS ACCESSION NUMBER: 1984:571174 CAPLUS

DOCUMENT NUMBER:

101:171174

TITLE: 2-Aryl-5, 5-dimethyl-1, 2, 4-triazolidin-3-one

derivatives

AUTHOR(S): Schantl, J.; Hebeisen, P.

CORPORATE SOURCE: Inst. Org. Pharm. Chem., Univ. Innsbruck, Innsbruck,

A-6020, Austria

SOURCE: Sci. Pharm. (1983), 51(4), 379-90

CODEN: SCPHA4; ISSN: 0036-8709

DOCUMENT TYPE: Journal LANGUAGE: German

AB RnC6H5-nNHN:CMe2 [Rn = H, 4-Cl, 3,4-Cl2, 4-Me(CH2)50, 4-02N] reacted with KZCN (Z = 0, S) in AcOH to give the corresponding triazolidinones I (Z = 0) or -thiones I (Z = S). Although I (Z = S) have antiinflammatory and analgesic properties I (Z = 0) had no noteworthy activity.

RnC6H5-nN:NCMe2N:C:Z, the acyclic oxidn. products of I, can be used for further syntheses. H2NCN was added to 4-ClC6H4NHN:CMe2.HCl to give

iminotriazolidine II which on oxidative ring cleavage gave
4-ClC6H4N:NCMe2NHCN.

IT 91027-32-6P

RN 91027-32-6 CAPLUS

$$\begin{array}{c} O \\ \parallel \\ \square H - C - NH - CH_2 - CH_2 - NMe_2 \end{array}$$

$$\begin{array}{c} N = N - C - Me \\ \downarrow \\ Me \end{array}$$

L37 ANSWER 12 OF 26 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 1983:615559 CAPLUS

DOCUMENT NUMBER: 99:215559

TITLE: Demulsification of bitumen emulsions using ionenes

INVENTOR(S): McCoy, David R.; McEntire, Edward E.

PATENT ASSIGNEE(S): Texaco Inc., USA

SOURCE: U.

U.S., 4 pp. CODEN: USXXAM

CODEM: OSKAP

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. HTM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE
US 4404096 A 19830913 US 1981-326459 19811202

AB Ionomer demulsifiers for breaking of tar-sand oil-in-water emulsions are prepd. rom ditertiarydiamines and dichlorohydrocarbons and have mol. wt.

>2600 / referably >10,000). Active compds. include

Me2NCH2CHLHT .-ClCH2-p-

PATENT INLE

TION:

C6H4CH2Cl copolymer [30619-25-1], Me2N(CH2)3NMe2-trans-C1CH2CH:CHCH2Cl copolymer [52193-09-6], 1,4-diazabicyclo[2,2,2]octane-trans-1,4-dichloro-2-butene copolymer [87836-94-0], and N,N,N',N'-tetramethyl-Jeffamine D-230-.alpha.,.alpha.'-dichloro-p-xylene copolymer [87935-68-0]. ΙT 69419-41-6 RL: USE: (Uses) (demulsifiers, for breaking of tar-sand emulsions) RN 69419-41-6 CAPLUS Urea, N,N'-bis[3-(dimethylamino)propyl]-, polymer with 1,4-bis(chloromethyl)benzene (9CI) (CA INDEX NAME) CM 1 CRN 50138-87-1 CMF CL H26 H4 O ($Me_2N - (CH_2)_3 - NH - C - NH - (CH_2)_3 - NMe_2$ CM Ź CRN 623-25-0 CMF 03 H8 C12 CH2Cl ClCH₂ L37 ANSWER 13 OF 26 CAPLUS COPYRIGHT 2001 ACS ACCESSION HTT BER: 1983:71669 CAPLUS DOCUMENT NAME ER: 98:71669 TITLE: $1\hbox{-}{\tt Phenoxy-3-ureidoalkylpropanolamine}\ \ {\tt derivatives}\ \ {\tt and}$ pharmaceutical compositions containing them PATENT ASSIGNEE(S): Imperial Chemical Industries PLC, UK SOURCE: Israeli, 19 pp. Addn. to Israeli 43,795. CODEN: ISXXAQ DOCUMENT TYDE: Patent **LANGU**AGE: English FAMILY ACT. WH. COUNT:

PATERT FO.	KIND	DATE		APPLICATION NO.	DATE			
IL 550:11	A1	19820531		IL 1978-55829	19781031			
PRIORITY ALDIM. INFO.	:		IL	1973-43795	19731210			
			GB	1977-52969	19771220			

Ý

AB

ΙT

84567-

-4P

```
Alkanc .mines I (R2 = H, 2-cyano, 2-Cl, 2-, 3-, 4-F, 2-Me, 2-MeO, 4-OH;
     ZR1 = JH2Ph, CHMePh; R2 = H, 4-OH, ZR1 = CH2CH2OH, CMe2CH2OH) and their
    acid addn. salts, having .beta.-adrenergic blocking activity (no data),
    were prepd. Treating PhO2CCl and K2CO3 in dioxane with PhCH2NH2 and
     stirring at room temp. 72 h gave PhCH2NHCO2Ph which was stirred with
     (H2NCH1)2 at room temp. 16 h to give PhCH2NHCONHCH2CH2NH2. This was
    heated with 1-(2-cyanophenoxy)-2,3-epoxypropane, H2O, and EtOH at
     90.deg: e. 16 h to give I (R2 = 2-cyano, ZR1 = CH2Ph).
     71676-1 -4P
IT
     RL: RC (Reactant); SPN (Synthetic preparation); PREP (Preparation)
        (propn. and aminolysis by, of epoxypropane deriv.)
RN
     71676-11-4 CAPLUS
     Urea, .:-(2-aminoethyl)-N'-(phenylmethyl)- (9CI) (CA INDEX NAME)
CN
           C_{i}
Ph-CH<sub>2</sub>-NII I-NH-CH<sub>2</sub>-CH<sub>2</sub>-NH<sub>2</sub>
L37 ANSWER 14 OF 26 CAPLUS COPYRIGHT 2001 ACS
ACCESSION TO MBER:
                         1983:71187 CAPLUS
                         98:71187
DOCUMENT HOW ER:
                         Direct spectrophotometric observation of an
TITLE:
                         O-acylisourea intermediate: concerted general acid
                         catalysis in the reaction of acetate ion with a
                         water-soluble carbodiimide
                         Ibrahim, Ibrahim T.; Williams, Andrew
AUTHOR(S):
                         Chem. Lab., Univ. Kent, Canterbury, CT2 7NZ, UK
CORPORATE 5 WRCE:
                         J. Chem. Soc., Perkin Trans. 2 (1982), (11), 1459-66
SOURCE:
                         CODEN: JCPKBH; ISSN: 0300-9580
DOCUMENT 1 . . :
                         Journal
                         English
LANGUAGE:
     Rate consts. for the formation and decompn. of intermediate
O-acylisous sas
     from cabodimide and carboxylic acids were measured in aq. media.
     O-acet in isourea from AcO- and
N-ethyl-N'
             - (trimethylammonio)propyl]carbo
             (I) has an acidic group of pK 6.8, and decomps. in its acid form
     diimi:
             ication by reaction with AcO- or H2O. Reaction of the
     as the
carboxylat
     anion to th I is general-acid catalyzed, and the D2O solvent isotope
effect
     indicates a rate-detg. proton transfer except for the oxonium ion acting
     as act. A mechanism involving proton transfer concerted with
             ilic attack by AcO- is consistent with the weak basicity of the
     nucle
             adduct. The 3rd-order term involving HOAc, AcO- and carbodiimide
     isou:
             .apprm.60% of the total reaction flux at pH 6.80 and 1 M total
             Ler concn. At this pH .apprx.40% of the reaction flux proceeds
     HOAc
             pwise mechanism with specific acid catalysis. Intramol.
     via a
             .cid catalysis occurs in the reaction of HO2CCEt2CO2- with I, and
     gene:
             ctive molarity compared with intermol. catalysis is 15 M. Attack
     the \epsilon
             Mylat - anions on I with N-(chloroethyl)morpholinium ion as the
     of c
     gene:
             acid .as a Broensted-type .beta.N of 0.46.
```

BEST AVAILABLE COPY

09/350,193

RL: : (Synthetic preparation); PREP (Preparation) a. of)(; RN 84567 -4 CAPLUS Urea, . [3-(dimethylamino)propyl]-N'-(2-phenylethyl)- (9CI) (CA INDEX CN NAME) 0|| Me2N- (CH: HH-C-NH-CH2-CH2-Ph L37 ANSWE: 5 OF 26 CAPLUS COPYRIGHT 2001 ACS ACCESSION BER: 1982:456677 CAPLUS DOCUMENT | ER: 97:56677 Cationic adsorbent and its use in removing anionic TITLE: products from aqueous solutions INVENTOR (Haase, Jaroslav; Palmberg, Roger Ciba-Geigy A.-G., Switz. PATENT AS. E(S): Eur. Pat. Appl., 31 pp. SOURCE: CODEN: EPXXDW DOCUMENT :: Patent LANGUAGE: German FAMILY ACC. UM. COUNT: 1 PATENT IN: :ATION: APPLICATION NO. DATE PATE: TO. KIND DATE ____ _____ EP 51 A2 19820505 EP 1981-810420 19811022 EP 5" АЗ 19820804 B1 19850502 IT, BE, CH, DE, FR, GB, IT, NL, SE US 4 + 294 A 19840103 US 1981-313061 19811019 AT 1981-810420 19811022 AT 13. Ε 19850515 ~3 DE 31 A1 19820527 DE 1981-3142153 19811023 .) A 19820429 FI 1981-3350 19811026 FΙ ٤ FI 7 FI 7... В 19860626 С 19861006 IL 1981-64112 19811026
CA 1981-388763 19811026
DK 1981-4733 19811027
BR 1981-6927 19811027 A1 19840930 A1 19850813 A 19820429 A 19820713 A 19820929 A1 19830201 IL 6 12 CA 1 3 DK ε . . 7 BR & 18 ZA 8 ZA 1981-7428 19811027 19830201 198**50923** ES 1981-506596 ES . 19811027 A3 2 23**1** SU 1981-3394405 19811027 SU 1 19820625 JP 🖯 231 A2 JP 1981-171541 19811028 JP (295 В4 19901024 А US 1983-520377 19830804 US 4 24 19841023 '. INFO.: PRIORITY . CH 1980-8016 19801028 US 1981-313061 19811019 EP 1981-810420 19811022 ion of an aminoplast precondensate with a compd. contg. amino AΒ The and

thiourea) groups, such as Me2N(CH2)3NHCONRCH2OH (I) (R = H or MeCl-quaternized I, gives cationic resins which are useful as

INVENTOR (S

PATENT ASS: . TE(S):

```
ts for anionic compds., e.g, for removing acid dyes from
             \cdotr. Thus, 51.7 g 40% soln. (pH 4.6) of I (R = H) was mixed with
             HOCH2NH)2CO and 10 mL 15% H2NSO3H, reflux for 90 min, evapd.,
     26.4
             tt 85-90.degree. for 15 h, pulverized, and added to water. The
     polym
             32539-88-6] particles were sepd. and dried to prep. 24.0 g
     resir
             t contg. 25.34% N.
     adso:
ΙT
     8253.
             ~3P
     RL: L
              (Preparation)
        (:
            on. of, as adsorbents for anionic materials)
     8253
RN
             -3 CAPLUS
CN
     Urea,
             3-(dimethylamino)propyl]-N'-(hydroxymethyl)-, polymer with
             yde and 1,3,5-triazine-2,4,6-triamine (9CI) (CA INDEX NAME)
     forma
     CM
             :9-68-2
     CRN
             117 N3 02
     CMF
                                                BEST AVAILABLE COPY
HO-CH2-N
           · NH- (CH2) 3- NMe2
     CM
     CRN
              78-1
     CMF
              5 N6
             .:2
     CM
              0 - 0
     CRN
     CMF
H_2C = 0
L37 ANSWE
             OF 26 CAPLUS COPYRIGHT 2001 ACS
                         1981:592212 CAPLUS
ACCESSION
              ER:
                         95:192212
DOCUMENT No.
                         Compositions for permanent waving of hair
TITLE:
```

Grollier, Jean Francois; Fourcadier, Chantal

Oreal S. A., Fr.

09/350,193 SOURCE: Fr. Demande, 18 pp. CODEN: FRXXBL DOCUMENT TYP :: Patent LANGUAGE: French FAMILY ACC. 1 M. COUNT: 3 PATENT INFORTATION: PATENT HO. KIND DATE APPLICATION NO. ____ _____ FR 2465 13 A2 19810327 FR 1979-30586 B2 FR 2465-18 19830610 BE 876 A1 BE 1979-195741 19791214 CH 6404 A 19840113 CH 1979-5592 US 434*. 2 A 19820907 US 1980-158271 US 4571.32 US 4970166 A 19860401 US 1982-406036 Α 19901113 US 1988-235955 PRIORITY APP : INFO.: US 1979-48585 BE 1979-195741 CH 1979-5592 IT 1979-68281 FR 1978-17899 CA 1979-329838 DE 1979-2924230 GB 1979-20878 JP 1979-75560

US 1986-845245 19860328

Quaternary ammonium polymers, e.g. N,N'-bis(3-dimethylaminopropyl)
urea-limated a.,.beta.-dichloroethyl ether copolymer (I) [68555-36-2], were prepd. It used in compns. for permanent waving of hair. Thus, I was prepd. It refluxing 0.2 mol of each of the corresponding monomers. Hair was trained with a compn. contg. thioglycolic acid 8, NH40H q.s.p. pH 7, NH4HCO3 1.4, dimethyldistearylammonium chloride (II) 0.2, I 3, oxyethy nated oleyl alc. 1, perfume, and water q.s.p. 100 g. After 5-15 min, the hair was rinsed, and treated with an oxidizing compn. contg. II 0.3, phacetin 0.1, citric acid 0.3, oxyethylenated nonylphenol 1, color and permanent waving of hair was easy to comb.

FR 1979-30586

US 1980-158271

DATE

19791213

19790614

19790614

19800610

19820806

19880823

19790613

19790614

19790614

19790614

19780615

19790615

19790615

19790615

19790615

19791213

19800610

IT 70698- 9P

RL: Pl: (Preparation)

(p: .. of, for hair wave-setting compns.)

RN 70698- 9 CAPLUS

CN Urea, : '-bis[3-{dimethylamino}propyl]-, polymer with 1,4-bi promomethyl)benzene (9CI) (CA INDEX NAME)

CM 1

CRN 5 8-87-1 CMF C: H26 N4 0

2 CM

CRN 623-24-5 CMF C8 H8 Br2

CH2Br BrCH₂

L37 ANSWER 17 OF 26 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

1981:121140 CAPLUS

DOCUMENT NUMBER:

94:121140

TITLE:

Alkanolamine derivatives with .beta.-adrenergic

blocking activity

INVENTOR(S):

Smith, Leslie Harold

PATENT ASSIGNEE(S):

Imperial Chemical Industries Ltd., Engl.

SOURCE:

Brit., 9 pp. Addn. to Brit. 1,455,116.

CODEN: BRXXAA

DOCUMENT TYPE:

Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE -----19800820 GB 1573359 A GB 1978-52969 19780518

The alkanolamine derivs., ROCH2CH(OH)CH2NHZ1NHCONHZ2R1 (R = R1 = aryl, Z1 = Z2 = ...lkylene; R = H, R1 = aryl, Z1 = alkylene, Z2 = alkyleneoxy) were prepd. as .beta.-adrenergic blockers with cardioselective action. E.g., reaction of 1-(2-cyanophenoxy)-2,3-epoxypropane with

1-(.beta.-aminoethyl)-

3-benzylurea gave

1-(2-cyanophenoxy)-3-.beta.-(3-benzylureidoethyl)amino-2-

propanol. For treatment of heart disease the active compd. was used at

or

oral dome of 20-600 mg daily or 1-20 mg i.v. and for treatment of acute

chronic heart failure the dose was 10-200 mg orally or 1-100 mg i.v.

IT 71676-11-4P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)

(prepa. and reaction of, with epoxypropane deriv.)

RN 71676-11-4 CAPLUS

CN Urea, N-(2-aminoethyl)-N'-(phenylmethyl)- (9CI) (CA INDEX NAME)

L37 ANSWER 18 OF 26 CAPLUS COPYRIGHT 2001 ACS ACCESSION NUMBER: 1981:57984 CAPLUS DOCUMENT NUMBER: 94:57984 TITLE: Potential inhibitors of nucleotide biosynthesis. 1. Nitrosoureidonucleosides. 2 Montgomery, John A.; Thomas, H. Jeanette; Brockman, AUTHOR(S): R. Wallace; Wheeler, Glynn P. CORPORATE SOURCE: Kettering-Meyer Lab., South. Res. Inst., Birmingham, AL, 35255, USA SOURCE: J. Med. Chem. (1981), 24(2), 184-9CODEN: JMCMAR; ISSN: 0022-2623 DOCUMENT TYPE: Journal LANGUAGE: English AΒ The title compds. I (R = H, Me, or cyclohexyl; R1 and R2 = H or NO; R3 = hypoxanthin-9-yl, thymin-1-yl, or uracil-1-yl; R4 = H or OH) were prepd. and evaluated for alkylating activity. The low level of biol. activity of. I is apparently due to their stability compared to the known nitrosourea compds. ΙT 75930-38-0P RL: SPM (Synthetic preparation); PREP (Preparation) (prepn. of) RN 75930-39-0 CAPLUS CN Urea, N-(2-aminoethyl)-N'-methyl-, compd. with 2,4,6-trinitrophenol (1:1) (9CI) (:A INDEX NAME) CM 1 CRN 75:30-29-9 CMF C4 H11 N3 O 0 MeNH-C-NH-CH2-CH2-NH2

CM

CRN 89-99-1 CMF Ct 113 N3 07

2

FILE 'USPATFULL' ENTERED AT 11:03:35 ON 01 JUN 2001
CA INDEXING COPYRIGHT (C) 2001 AMERICAN CHEMICAL SOCIETY (ACS)

FILE COVERS 1971 TO PATENT PUBLICATION DATE: 29 May 2001 (20010529/PD)
FILE LAST UPDATED: 29 May 2001 (20010529/ED)
HIGHEST PATENT NUMBER: US8411134
CA INDEXING IS CURRENT THROUGH 29 May 2001 (20010529/UPCA)
ISSUE CLASS FIELDS (/INCL) CURRENT THROUGH: 29 May 2001 (20010529/PD)
REVISED CLASS FIELDS (/NCL) LAST RELOADED: Apr 2001
USPTO MANUAL OF CLASSIFICATIONS THESAURUS ISSUE DATE: Apr 2001

>>> Page images are available for patents from 1/1/1997. Current <<< >>> week patent text is typically loaded by Thursday morning and <<< >>> page images are available for display by the end of the day. <<< >>> Image data for the /FA field are available the following week. <<

>>> Complete CA file indexing for chemical patents (or equivalents) <<< >>> is included in file records. A thesaurus is available for the >>> USPTO Manual of Classifications in the /NCL, /INCL, and /RPCL <<< >>> fields. This thesaurus includes catchword terms from the <<< >>> USPTO/MOC subject headings and subheadings. Thesauri are also <<< >>> available for the WIPO International Patent Classification <<< >>> (IPC) Manuals, editions 1-6, in the /IC1, /IC2, /IC3, /IC4, <<< >>> /IC5, and /IC (/IC6) fields, respectively. The thesauri in <<< >>> the /IC5 and /IC fields include the corresponding catchword <<< >>> terms from the IPC subject headings and subheadings. <<<

This file contains CAS Registry Numbers for easy and accurate substance identification.

=> s 122 L25 7 L22

=> d ibib ab hitstr 1-7

L25 ANSWER 1 OF 7 USPATFULL

ACCESSION NUMBER: 94:80134 USPATFULL

TITLE: 3-aminopropoxyphenyl derivatives, their preparation

and

pharmaceutical compositions containing them

INVENTOR(S): Berthold, Richard, 9 Ahornstrasse, CH-4103 Bottmingen,

Switzerland

Louis, William J., 3 Balmoral Avenue, Kew, 3101

Victoria, Australia

PATENT INFORMATION: US 5347050 19940913
APPLICATION INFO.: US 1993-46937 19930413 (8)
RELATED APPLN. INFO.: Continuation of Ser. No. US 1991-782791, filed on 21 Oct 1991, now abandoned which is a continuation of Ser.

No. US 1990-584306, filed on 17 Sep 1990, now abandoned which is a continuation of Ser. No. US 1990-474185,

filed on 2 Feb 1990, now abandoned which is a

continuation of Ser. No. US 1989-399721, filed on 25 Aug 1989, now abandoned which is a continuation of

Ser.

No. US 1989-307028, filed on 3 Feb 1989, now abandoned which is a continuation of Ser. No. US 1988-173845, filed on 28 Mar 1988, now abandoned which is a continuation of Ser. No. US 1986-897557, filed on 18 Aug 1986, now abandoned which is a continuation of

Ser.

No. US 1985-778831, filed on 23 Sep 1985, now

abandoned

which is a continuation of Ser. No. US 1984-567471, filed on 3 Jan 1984, now abandoned which is a division of Ser. No. US 1981-318292, filed on 4 Nov 1981, now

patented, Pat. No. US 4425362

	NUMBER	DATE
PRIORITY INFORMATION:	CH 1980-8249	19801106
	СН 1980-9347	19801218
	CH 1981-4073	19810619
	CH 1991-407481	19910619
DOCUMENT TYPE:	Utility	
PRIMARY EXAMINER:	Dees, Jose G.	
	~ ' ~ '	

ASSISTANT EXAMINER:

Carr, Deborah D.

LEGAL REPRESENTATIVE: NUMBER OF CLAIMS:

Sughrue, Mion, Zinn, Macpeak & Seas

EXEMPLARY CLAIM: LINE COUNT: 1090

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The compounds of formula I, ##STR1## wherein the substituents have various significances, and physiologically acceptable hydrolyzable derivatives thereof having the hydroxy group in the 2 position of the 3-aminopropoxy side chain in esterified form, are useful as cardioselective .beta.-adrenoceptor blocking agents.

ΙT 75930-29-9

(reaction of, with (epoxypropoxy)benzene derivs.)

RN 75930-29-9 USPATFULL

Urea, N-(2-aminoethyl)-N'-methyl- (9CI) (CA INDEX NAME) CN

0 MeNH-C-NH-CH2-CH2-NH2

L25 ANSWER 2 OF 7 USPATFULL

90:48806 USPATFULL ACCESSION NUMBER:

TITLE: 2-hydroxypropylamine aryl ester derivatives and

pharmaceutical use

INVENTOR(S): Kam, Sheung T., Vernon Hills, IL, United States

Matier, William L., Libertyville, IL, United States

PATENT ASSIGNEE(S): E. I. Du Pont de Nemours and Company, Wilmington, DE,

United States (U.S. corporation)

PATENT ASSIGNEE(S):

NUMBER DATE US 4935421 19900619 PATENT INFORMATION: US 1989-318147 19890301 (7) APPLICATION INFO .: RELATED APPLN. INFO.: Division of Ser. No. US 1986-838082, filed on 10 Mar 1986, now patented, Pat. No. US 4810717 which is a division of Ser. No. US 1981-320773, filed on 21 Nov 1981, now patented, Pat. No. US 4582855 Utility DOCUMENT TYPE: Ramsuer, Robert W. PRIMARY EXAMINER: NUMBER OF CLAIMS: 27 EXEMPLARY CLAIM: 1 1470 LINE COUNT: CAS INDEXING IS AVAILABLE FOR THIS PATENT. Novel compounds of the general formula ##STR1## wherein Ar represents a substituted or unsubstituted aromatic or heterocyclic group; W represents alkylene of from 1 to about 10 carbon atoms; and B represents --NR.sub.2 COR.sub.1, --NR.sub.2 CONR.sub.1 R.sub.3, --NR.sub.2 SO.sub.2 R.sub.1, --NR.sub.2 SO.sub.2 NR.sub.1 R.sub.3, or --NR.sub.2 COOR.sub.1 wherein R.sub.1, R.sub.2 and R.sub.3 may be the same or different and may be hydrogen, alkyl, alkoxyalkyl, cycloalkyl, alkenyl, alkynyl, aryl, heteroaryl, or aralkyl, except that R.sub.1 is not hydrogen when B is --NR.sub.2 SO.sub.2 R.sub.1 or --NR.sub.2 COOR.sub.1, or R.sub.1 and R.sub.3 may together with N form a 5 to 7 membered heterocyclic group; and the pharmaceutically acceptable salts thereof. These compounds exhibit .beta.-adrenergic blocking activity and are also useful in the treatment of glaucoma. ΙT 75930-29-9P (prepn. of) RN 75930-29-9 USPATFULL Urea, N-(2-aminoethyl)-N'-methyl- (9CI) (CA INDEX NAME) CN MeNH-C-NH-CH2-CH2-NH2 L25 ANSWER 3 OF 7 USPATFULL ACCESSION NUMBER: 89:17318 USPATFULL 2-hydroxypropylamine aryl ester derivatives TITLE: Kam, Sheung T., Vernon Hills, IL, United States INVENTOR(S):

	NOMBER	DATE	
PATENT INFORMATION:	US 4810717	19890307	
APPLICATION INFO.:	US 1986-838082	19860310 (6)	
RELATED APPLN. INFO.:	Division of Ser.	No. US 1981-320773	, filed on 12 Nov
	1981, now patent	ed, Pat. No. US 458	2855

MILIMPED

United States (U.S. corporation)

DAME

Matier, William L., Libertyville, IL, United States E. I. du Pont de Nemours and Company, Wilmington, DE,

DOCUMENT TYPE:

Utility

PRIMARY EXAMINER:

Lee, Mary C.

ASSISTANT EXAMINER:

Whittenbaugh, Robert C.

LEGAL REPRESENTATIVE:

Fato, Gildo E.

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

33 1,11

LINE COUNT:

1764

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Novel compounds of the general formula ##STR1## wherein Ar represents a substituted or unsubstituted aromatic or heterocyclic group; W represents alkylene of from 1 to about 10 carbon atoms; and B

represents

--NR.sub.2 COR.sub.1, --NR.sub.2 CONR.sub.1 R.sub.3, --NR.sub.2

SO.sub.2

R.sub.1, --NR.sub.2 SO.sub.2 NR.sub.1 R.sub.3, or --NR.sub.2 COOR.sub.1 wherein R.sub.1, R.sub.2 and R.sub.3 may be the same or different and may be hydrogen, alkyl, alkoxyalkyl, cycloalkyl, alkenyl, alkynyl,

aryl,

heteroaryl, or aralkyl, except that R.sub.1 is not hydrogen when B is --NR.sub.2 SO.sub.2 R.sub.1 or --NR.sub.2 COOR.sub.1, or R.sub.1 and R.sub.3 may together with N form a 5 to 7 membered heterocyclic group; and the pharmaceutically acceptable salts thereof. These compounds exhibit .beta.-adrenergic blocking activity and are also useful in the treatment of glaucoma.

IT 75930-29-9P

(prepn. of)

RN 75930-29-9 USPATFULL

Urea, N-(2-aminoethyl)-N'-methyl- (9CI) (CA INDEX NAME) CN

MeNH-C-NH-CH2-CH2-NH2

L25 ANSWER 4 OF 7 USPATFULL

ACCESSION NUMBER:

89:4612 USPATFULL

TITLE:

INVENTOR(S):

2-hydroxypropylamine heteroaryl ester derivatives

Kam, Sheung T., Chicago, IL, United States

Matier, William L., Libertyville, IL, United States Patil, Ghanshyam, Vernon Hills, IL, United States

Mai, Khuong H. X., Waukegan, IL, United States

E. I. Du Pont de Nemours and Company, Wilmington, DE, PATENT ASSIGNEE(S):

United States (U.S. corporation)

NUMBER DATE _____

PATENT INFORMATION:

US 4798892 19890117

US 1986-851629 19860414 (6)

APPLICATION INFO.: RELATED APPLN. INFO.:

Continuation-in-part of Ser. No. US 1981-320773, filed

on 12 Nov 1981, now patented, Pat. No. US 4582855,

issued on 15 Apr 1986

DOCUMENT TYPE:

Utility

PRIMARY EXAMINER:

Raymond, Richard L.

LEGAL REPRESENTATIVE: Fato, Gildo E.

NUMBER OF CLAIMS: 9
EXEMPLARY CLAIM: 1
LINE COUNT: 1391

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention relates to compounds of the general formula ##STR1## wherein Ar represents a substituted or unsubstituted heterocyclic group; W represents alkylene of from 1 to about 10 carbon atoms; and B represents --NR.sub.2 COR.sub.1, --NR.sub.2 CONR.sub.1 R.sub.3, --NR.sub.2 SO.sub.2 R.sub.1, NR.sub.2 SO.sub.2 NR.sub.1 R.sub.3, or --NR.sub.2 COOR.sub.1, wherein R.sub.1, R.sub.2 and R.sub.3 may be alike or different and may be hydrogen, alkyl, alkoxyalkyl cycloalkyl, alkenyl, alkynyl, aryl, heteroaryl, or aralkyl, except that R.sub.1 is not hydrogen when B is --NR.sub.2 SO.sub.2 R.sub.1 or --NR.sub.2 COOR.sub.1, or R.sub.1 and R.sub.3 may together with N form

5 to 7 membered heterocyclic group and the pharmaceutically acceptable salts thereof. The compounds exhibit beta-adrenergic blocking activity and are also useful in the treatment of glaucoma.

IT 122036-80-0P

(prepn. and reaction of, in prepn. of .beta.-adrenergic blockers)

RN 122036-80-0 USPATFULL

CN Urea, N-(2-aminoethyl)-N'-methyl-, monohydrochloride (9CI) (CA INDEX NAME)

, O || MeNH-C-NH-CH₂-CH₂-NH₂

HCl

L25 ANSWER 5 OF 7 USPATFULL

ACCESSION NUMBER: 87:3253 USPATFULL

TITLE: Para-substituted 3-phenoxy-1-carbonylamino-alkylamino-

propanol compounds, beta receptor blocking

compositions

and use

INVENTOR(S): Gustafsson, Bill B. R., Bollebygd, Sweden

Hedberg, Sven A., Gr.ang.bo, Sweden Lundgren, Bo T., Frilles.ang.s, Sweden

PATENT ASSIGNEE(S): Aktiebolaget Hassle, Molndal, Sweden (non-U.S.

corporation)

NUMBER DATE

PATENT INFORMATION: US 4636501 19870113

APPLICATION INFO.: US 1985-757763 19850722 (6)

RELATED APPLN. INFO.: Continuation of Ser. No. US 1984-621147, filed on 18

Jun 1984, now abandoned which is a continuation of

Ser.
No. US 1983-482266, filed on 5 Apr 1983, now abandoned

which is a continuation-in-part of Ser. No. US

aryl,

1982-450006, filed on 15 Dec 1982, now abandoned

NUMBER DATE SE 1981-7574 19811217 PRIORITY INFORMATION: Utility DOCUMENT TYPE: Ramsuer, Robert W. PRIMARY EXAMINER: Brumbaugh, Graves, Donohue & Raymond LEGAL REPRESENTATIVE: NUMBER OF CLAIMS: 1,9 EXEMPLARY CLAIM: 1017 LINE COUNT: CAS INDEXING IS AVAILABLE FOR THIS PATENT. Compounds of the formula ##STR1## having beta receptor blocking properties, are disclosed. IT 75930-29-9 (ring cleavage by, of glycidyl aryl ethers) RN 75930-29-9 USPATFULL Urea, N-(2-aminoethyl)-N'-methyl- (9CI) (CA INDEX NAME) CN MeNH-C-NH-CH2-CH2-NH2 L25 ANSWER 6 OF 7 USPATFULL 86:21877 USPATFULL ACCESSION NUMBER: Aromatic and esters of hydroxypropylamines TITLE: Kam, Sheung T., Vernon Hills, IL, United States INVENTOR(S): Matier, William L., Libertyville, IL, United States American Hospital Supply Corporation, Evanston, IL, PATENT ASSIGNEE(S): United States (U.S. corporation) DATE NUMBER _____ US 4582855 19860415 PATENT INFORMATION: APPLICATION INFO.: US 1981-320773 19811112 (6) DOCUMENT TYPE: Utility PRIMARY EXAMINER: Jiles, Henry R. Whittenbaugh, Robert C. ASSISTANT EXAMINER: Kanady, Mary Jo; Barbeau, Donald L.; Fato, Gildo E. LEGAL REPRESENTATIVE: 57 NUMBER OF CLAIMS: 1,30 EXEMPLARY CLAIM: 1804 LINE COUNT: CAS INDEXING IS AVAILABLE FOR THIS PATENT. Novel compounds of the general formula ##STR1## wherein Ar represents a AΒ substituted or unsubstituted aromatic or heterocyclic group; W represents alkylene of from 1 to about 10 carbon atoms; and B represents --NR.sub.2 COR.sub.1, --NR.sub.2 CONR.sub.1 R.sub.3, --NR.sub.2 SO.sub.2 R.sub.1, --NR.sub.2 SO.sub.2 NR.sub.1 R.sub.3, or --NR.sub.2 COOR.sub.1

wherein R.sub.1, R.sub.2 and R.sub.3 may be the same or different and may be hydrogen, alkyl, alkoxyalkyl, cycloalkyl, alkenyl, alkynyl,

heteroaryl, or aralkyl, except that R.sub.1 is not hydrogen when B is --NR.sub.2 SO.sub.2 R.sub.1 or --NR.sub.2 COOR.sub.1, or R.sub.1 and R.sub.3 may together with N form a 5 to 7 momoered heterocyclic group; and the pharmaceutically acceptable salts thereof. These compounds exhibit .beta.-adrenergic blocking activity and are also useful in the treatment of glaucoma.

IT 75930-29-9P

(prepn. of)

RN 75930-29-9 USPATFULL

CN Urea, N-(2-aminoethyl)-N'-methyl- (9CI) (CA INDEX NAME)

O || MeNH-C-NH-CH2-CH2-NH2

L25 ANSWER 7 OF 7 USPATFULL

ACCESSION NUMBER: 84:2057 USPATFULL

TITLE: 3-Aminopropoxyphenyl derivatives and pharmaceutical

compositions containing them

INVENTOR(S): Berthold, Richard, Bottmingen, Switzerland

Louis, William J., Kew, Australia

PATENT ASSIGNEE(S): Sandoz Ltd., Basel, Switzerland (non-U.S. corporation)

DOCUMENT TYPE: Utility

PRIMARY EXAMINER: Torrence, Dolph H.

LEGAL REPRESENTATIVE: Sharkin, Gerald D.; Honor, Robert S.

NUMBER OF CLAIMS: 11 EXEMPLARY CLAIM: 1,10 LINE COUNT: 1101

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB The compounds of formula I, ##STR1## wherein the substituents have various significances, and physiologically acceptable hydrolyzable derivatives thereof having the hydroxy group in the 2 position of the 3-aminopropoxy side chain in esterified form, are useful as cardioselective .beta.-adrenoceptor blocking agents.

IT 75930-29-9

(reaction of, with (epoxypropoxy)benzene derivs.)

RN 75930-29-9 USPATFULL

CN Urea, N-(2-aminoethyl)-N'-methyl- (9CI) (CA INDEX NAME)

O || MeNH-C-NH-CH₂-CH₂-NH₂

=> file reg SINCE FILE COST IN U.S. DOLLARS TOTAL ENTRY SESSION FULL ESTIMATED COST 36.50 635.00 DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) SINCE FILE TOTAL ENTRY SESSION CA SUBSCRIBER PRICE 0.00 -15.88

FILE 'REGISTRY' ENTERED AT 11:05:41 ON 01 JUN 2001 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS.
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STRUCTURE FILE UPDATES: 30 MAY 2001 HIGHEST RN 339046-06-9 DICTIONARY FILE UPDATES: 30 MAY 2001 HIGHEST RN 339046-06-9

TSCA INFORMATION NOW CURRENT THROUGH January 11, 2001

Please note that search-term pricing does apply when conducting SmartSELECT searches.

Structure search limits have been increased. See HELP SLIMIT for details.

=> d his

(FILE 'HOME' ENTERED AT 10:38:39 ON 01 JUN 2001)

FILE 'REGISTRY' ENTERED AT 10:38:44 ON 01 JUN 2001 L1STRUCTURE UPLOADED L2 50 S L1 L3 36270 S L1 FULL STRUCTURE UPLOADED L414060 S L4 FULL SUB=L3 L5 5399 S L5 AND 3/N L6 734 S ETHYL(L) DIMETHYL(L) AMINO(L) PROPYL(L) UREA L70 S UREA, "N-ETHYL-N'-(3-DIMETHYLAMINOPROPYL)-" L84 S ETHYL(L) DIMETHYLAMINOPROPYL(L) UREA L9 L103 S L9 AND 1/NC 1 S L10 AND 1/0 L11 FILE 'CAPLUS' ENTERED AT 10:52:57 ON 01 JUN 2001 15 S L11 L12

L13 2 S L11/THU

ETTE LUCRATEULL ENTERED AT 10.55.19 ON 01 TUN 200

FILE 'USPATFULL' ENTERED AT 10:55:19 ON 01 JUN 2001 L14 2 S L11

FILE 'MARPAT' ENTERED AT 10:56:49 ON 01 JUN 2001

L15 0 S L11 L16 0 S L9

FILE 'BEILSTEIN' ENTERED AT 10:57:15 ON 01 JUN 2001

L17 1 S L11

FILE 'REGISTRY' ENTERED AT 10:57:49 ON 01 JUN 2001

L18 STRUCTURE UPLOADED

L19 11247 S L18 FULL SUB=L3

L20 50 S L18

L21 0 S L18 CSS

L22 8 S L18 CSS FULL

FILE 'CAPLUS' ENTERED AT 11:01:43 ON 01 JUN 2001

L23 10 S L22

L24 0 S L22/THU

FILE 'USPATFULL' ENTERED AT 11:03:35 ON 01 JUN 2001

L25 7 \$ L22

FILE 'REGISTRY' ENTERED AT 11:05:41 ON 01 JUN 2001

=> s 118 full

FULL SEARCH INITIATED 11:05:52 FILE 'REGISTRY'
FULL SCREEN SEARCH COMPLETED - 54245 TO ITERATE

100.0% PROCESSED 54245 ITERATIONS

11247 ANSWERS

SEARCH TIME: 00.00.05

L26 11247 SEA SSS FUL L18

=> d scan

L26 11247 ANSWERS REGISTRY COPYRIGHT 2001 ACS

IN Urea,

N-[1-[(3,4-dichlorophenyl)methyl]-1,6-dihydro-6-oxo-3-pyridinyl]-N'[3-(dimethylamino)propyl]- (9CI)

MF C18 H22 C12 N4 O2

$$Me_{2}N-(CH_{2})_{3}-NH-C-NH$$

$$N-CH_{2}$$

$$C1$$

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):2

L26 11247 ANSWERS REGISTRY COPYRIGHT 2001 ACS

IN Urea, N-[2-[[[(4-methylphenyl)amino]carbonyl]amino]ethyl]-N'-(4-

nitrophenyl)-N-(4-phenoxyphenyl)- (9CI)
MF C29 H27 N5 O5

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

=> Uploading 489.str

L27 STRUCTURE UPLOADED

=> d his

(FILE 'HOME' ENTERED AT 10:38:39 ON 01 JUN 2001)

FILE 'REGISTRY' ENTERED AT 10:38:44 ON 01 JUN 2001 STRUCTURE UPLOADED L1L2 50 S L1 L3 36270 S L1 FULL L4STRUCTURE UPLOADED L5 14060 S L4 FULL SUB=L3 L6 5399 S L5 AND 3/N L7 734 S ETHYL(L) DIMETHYL(L) AMINO(L) PROPYL(L) UREA L8 0 S UREA, "N-ETHYL-N'-(3-DIMETHYLAMINOPROPYL)-" L9 4 S ETHYL(L)DIMETHYLAMINOPROPYL(L)UREA

```
3 S L9 AND 1/NC
L10
            1 S L10 AND 1/O
L11
    FILE 'CAPLUS' ENTERED AT 10:52:57 ON 01 JUN 2001
      15 S L11
L12
            2 S L11/THU
L13
   FILE 'USPATFULL' ENTERED AT 10:55:19 ON 01 JUN 2001
L14
             2 S L11
    FILE 'MARPAT' ENTERED AT 10:56:49 ON 01 JUN 2001
L15
             0 S L11
L16
             0 S L9
    FILE 'BEILSTEIN' ENTERED AT 10:57:15 ON 01 JUN 2001
L17
             1 S L11
   FILE 'REGISTRY' ENTERED AT 10:57:49 ON 01 JUN 2001
              STRUCTURE UPLOADED
L18
         11247 S L18 FULL SUB=L3
L19
            50 S L18
L20
             0 S L18 CSS
L21
             8 S L18 CSS FULL
L22
    FILE 'CAPLUS' ENTERED AT 11:01:43 ON 01 JUN 2001
L23
            10 S L22
             0 S L22/THU
L24
    FILE 'USPATFULL' ENTERED AT 11:03:35 ON 01 JUN 2001
L25
            7 S L22
    FILE 'REGISTRY' ENTERED AT 11:05:41 ON 01 JUN 2001
L26
        11247 S L18 FULL
               STRUCTURE UPLOADED
L27
=> s 127 sub=126 full
FULL SUBSET SEARCH INITIATED 11:08:23 FILE 'REGISTRY'
FULL SUBSET SCREEN SEARCH COMPLETED - 11247 TO ITERATE
100.0% PROCESSED 11247 ITERATIONS
                                                           3145 ANSWERS
SEARCH TIME: 00.00.02
        3145 SEA SUB=L26 SSS FUL L27
=> d scan
L28 3145 ANSWERS REGISTRY COPYRIGHT 2001 ACS
IN Urea,
N-[6-bromo-2-(3-thienyl)-4-quinolinyl]-N'-[3-(dimethylamino)propyl]-
    (9CI)
MF C19 H21 Br N4 O S
```

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):1

L28 3145 ANSWERS REGISTRY COPYRIGHT 2001 ACS
IN Urea, N-(2,3-dichlorophenyl)-N'-[6-[(4-nitrophenyl)amino]hexyl]- (9CI)
MF C19 H22 C12 N4 O3

$$NH - (CH2)6 - NH - C - NH - C1$$

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

=> Uploading 489.str

opioading 403.5ci

L29 STRUCTURE UPLOADED

=> d his

(FILE 'HOME' ENTERED AT 10:38:39 ON 01 JUN 2001)

FILE 'REGISTRY' ENTERED AT 10:38:44 ON 01 JUN 2001 L1STRUCTURE UPLOADED L2 50 S L1 36270 S L1 FULL L3 STRUCTURE UPLOADED L414060 S L4 FULL SUB=L3 L5 5399 S L5 AND 3/N L6 734 S ETHYL(L)DIMETHYL(L)AMINO(L)PROPYL(L)UREA L7 0 S UREA, "N-ETHYL-N'-(3-DIMETHYLAMINOPROPYL)-" rs4 S ETHYL(L) DIMETHYLAMINOPROPYL(L) UREA L9 3 S L9 AND 1/NC L10 1 S L10 AND 1/0 L11 FILE 'CAPLUS' ENTERED AT 10:52:57 ON 01 JUN 2001 L12 15 S L11 L13 2 S L11/THU

FILE 'USPATFULL' ENTERED AT 10:55:19 ON 01 JUN 2001 L14 2 S L11

FILE 'MARPAT' ENTERED AT 10:56:49 ON 01 JUN 2001 0 S L11 L15 0 S L9 L16 FILE 'BEILSTEIN' ENTERED AT 10:57:15 ON 01 JUN 2001 L17 1 S L11 FILE 'REGISTRY' ENTERED AT 10:57:49 ON 01 JUN 2001 STRUCTURE UPLOADED L18 11247 S L18 FULL SUB=L3 L19 L20 50 S L18 L21 0 S L18 CSS 8 S L18 CSS FULL L22 FILE 'CAPLUS' ENTERED AT 11:01:43 ON 01 JUN 2001 10 S L22 L23 0 S L22/THU L24 FILE 'USPATFULL' ENTERED AT 11:03:35 ON 01 JUN 2001 7 S L22 L25 FILE 'REGISTRY' ENTERED AT 11:05:41 ON 01 JUN 2001 L26 11247 S L18 FULL STRUCTURE UPLOADED L27 3145 S L27 FULL SUB=L26 L28 L29 STRUCTURE UPLOADED => s 129 sub=128 full FULL SUBSET SEARCH INITIATED 11:13:36 FILE 'REGISTRY' FULL SUBSET SCREEN SEARCH COMPLETED - 3145 TO ITERATE 1523 ANSWERS 100.0% PROCESSED 3145 ITERATIONS SEARCH TIME: 00.00.04 L30 1523 SEA SUB=L28 SSS FUL L29 => d scan

Absolute stereochemistry.

C22 H38 N4 O10

L30 1523 ANSWERS

MF

REGISTRY COPYRIGHT 2001 ACS

4,6,15,17-Tetraazaeicosane-1,3,18,20-tetracarboxylic acid, 5,16-dioxo-,

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

3,18-dimethyl ester, (3S,18S)- (9CI)

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=> d his
     (FILE 'HOME' ENTERED AT 10:38:39 ON 01 JUN 2001)
     FILE 'REGISTRY' ENTERED AT 10:38:44 ON 01 JUN 2001
                STRUCTURE UPLOADED
L1
L2
             50 S L1
L3
          36270 S L1 FULL
L4
                STRUCTURE UPLOADED
          14060 S L4 FULL SUB=L3
L5
           5399 S L5 AND 3/N
L6
            734 S ETHYL(L) DIMETHYL(L) AMINO(L) PROPYL(L) UREA
L7
              0 S UREA, "N-ETHYL-N'-(3-DIMETHYLAMINOPROPYL)-"
^{\text{L8}}
              4 S ETHYL(L) DIMETHYLAMINOPROPYL(L) UREA
L9
L10
              3 S L9 AND 1/NC
L11
              1 S L10 AND 1/0
     FILE 'CAPLUS' ENTERED AT 10:52:57 ON 01 JUN 2001
             15 S L11
L12
              2 S L11/THU
L13
     FILE 'USPATFULL' ENTERED AT 10:55:19 ON 01 JUN 2001
              2 S L11
L14
     FILE 'MARPAT' ENTERED AT 10:56:49 ON 01 JUN 2001
L15
              0 S L11
              0 S L9
L16
     FILE 'BEILSTEIN' ENTERED AT 10:57:15 ON 01 JUN 2001
L17
             1 S L11
     FILE 'REGISTRY' ENTERED AT 10:57:49 ON 01 JUN 2001
L18
               STRUCTURE UPLOADED
L19
          11247 S L18 FULL SUB=L3
             50 S L18
L20
              0 S L18 CSS
L21
L22
              8 S L18 CSS FULL
     FILE 'CAPLUS' ENTERED AT 11:01:43 ON 01 JUN 2001
L23
             10 S L22
L24
             0 S L22/THU
     FILE 'USPATFULL' ENTERED AT 11:03:35 ON 01 JUN 2001
L25
              7 S L22
     FILE 'REGISTRY' ENTERED AT 11:05:41 ON 01 JUN 2001
L26
          11247 S L18 FULL
L27
                STRUCTURE UPLOADED
L28
           3145 S L27 FULL SUB=L26
L29
                STRUCTURE UPLOADED
           1523 S L29 FULL SUB=L28
L30
=> s 130 and 1/o
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2846307 1/0

381 L30 AND 1/0

L31

09/350,193

=> d scan

L31 381 ANSWERS REGISTRY COPYRIGHT 2001 ACS

IN 1-Dodecanaminium, N, N-dimethyl-N-[2-[[[[3-[(3,3,4,4,5,5,6,6,6-

nonafluorohexyl)thio]propyl]amino]carbonyl]amino]ethyl]-, bromide (9CI)

MF C26 H47 F9 N3 O S . Br

● Br-

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

=> s 131 not s/els 4193743 S/ELS L32 331 L31 NOT S/ELS

=> d scan

L32 331 ANSWERS REGISTRY COPYRIGHT 2001 ACS
IN Urea, N-(4-aminobutyl)-N'-(triphenylmethyl)- (9CI)
MF C24 H27 N3 O

$$^{\rm O}_{||}$$
 Ph₃C-NH-C-NH-(CH₂)4-NH₂

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):1

L32 331 ANSWERS REGISTRY COPYRIGHT 2001 ACS IN Urea, N-[2-(dimethylamino)ethyl]-N'-ethyl-, monohydrochloride (9CI) MF C7 H17 N3 O . Cl H

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):1

L32 331 ANSWERS REGISTRY COPYRIGHT 2001 ACS

IN 1-Butanaminium, N,N,N-trimethyl-4-[[(methylamino)carbonyl]amino]- (9CI)

MF C9 H22 N3 O

CI COM

O || MeNH- C- NH- (CH2)4-N+Me3

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

=> file caplus COST IN U.S. DOLLARS SINCE FILE TOTAL. ENTRY SESSION FULL ESTIMATED COST 208.50 843.50 DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) SINCE FILE TOTAL ENTRY SESSION CA SUBSCRIBER PRICE 0.00 -15.88

FILE 'CAPLUS' ENTERED AT 11:14:57 ON 01 JUN 2001 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS.

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FILE COVERS 1947 - 1 Jun 2001 VOL 134 ISS 23 FILE LAST UPDATED: 30 May 2001 (20010530/ED)

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=> d his

(FILE 'HOME' ENTERED AT 10:38:39 ON 01 JUN 2001)

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FILE 'REGISTRY' ENTERED AT 10:38:44 ON 01 JUN 2001
L1
                 STRUCTURE UPLOADED
              50 S L1
L2
L3
           36270 S L1 FULL
L4
                 STRUCTURE UPLOADED
L5
           14060 S L4 FULL SUB=L3
L6
            5399 S L5 AND 3/N
L7
             734 S ETHYL(L) DIMETHYL(L) AMINO(L) PROPYL(L) UREA
               0 S UREA, "N-ETHYL-N'-(3-DIMETHYLAMINOPROPYL)-"
L8
L9
               4 S ETHYL(L)DIMETHYLAMINOPROPYL(L)UREA
L10
               3 S L9 AND 1/NC
L11
               1 S L10 AND 1/0
     FILE 'CAPLUS' ENTERED AT 10:52:57 ON 01 JUN 2001
L12
             15 S L11
L13
              2 S L11/THU
     FILE 'USPATFULL' ENTERED AT 10:55:19 ON 01 JUN 2001
L14
               2 S L11
     FILE 'MARPAT' ENTERED AT 10:56:49 ON 01 JUN 2001
L15
              0 S L11
L16
              0 S L9
     FILE 'BEILSTEIN' ENTERED AT 10:57:15 ON 01 JUN 2001
L17
              1 S L11
     FILE 'REGISTRY' ENTERED AT 10:57:49 ON 01 JUN 2001
L18
                STRUCTURE UPLOADED
L19
          11247 S L18 FULL SUB=L3
L20
             50 S L18
L21
              0 S L18 CSS
L22
              8 S L18 CSS FULL
     FILE 'CAPLUS' ENTERED AT 11:01:43 ON 01 JUN 2001
L23
             10 S L22
L24
              0 S L22/THU
     FILE 'USPATFULL' ENTERED AT 11:03:35 ON 01 JUN 2001
L25
              7 S L22
     FILE 'REGISTRY' ENTERED AT 11:05:41 ON 01 JUN 2001
L26
          11247 S L18 FULL
L27
                STRUCTURE UPLOADED
L28
           3145 S L27 FULL SUB=L26
L29
                STRUCTURE UPLOADED
L30
           1523 S L29 FULL SUB=L28
L31
           381 S L30 AND 1/0
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L32
              331 S L31 NOT S/ELS
      FILE 'CAPLUS' ENTERED AT 11:14:57 ON 01 JUN 2001
 => s 132/thu
             226 L32
         375285 THU/RL
 L33
             14 L32/THU
                   (L32 (L) THU/RL)
=> d ibib ab hitstr 1-14
L33 ANSWER 1 OF 14 CAPLUS COPYRIGHT 2001 ACS
ACCESSION NUMBER:
                            2001:338479 CAPLUS
TITLE:
                            Preparation of amides and ureas as activators of
                            soluble guanylate cyclase
INVENTOR(S):
                            Selwood, David; Glen, Robert; Reynolds, Karen;
                           Wishart, Grant
PATENT ASSIGNEE(S):
                           University College London, UK
SOURCE:
                           PCT Int. Appl., 101 pp.
                           CODEN: PIXXD2
DOCUMENT TYPE:
                           Patent
LANGUAGE:
                           English
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                      KIND DATE
      PATENT NO.
                                              APPLICATION NO. DATE
      -----
                       ----
                                               -----
     WO 2001032604
                              20010510
                       A1
                                              WO 2000-GB4249 20001106
         W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU,
              SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN,
              YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
          RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
              DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,
              BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
PRIORITY APPLN. INFO.:
                                        GB 1999-26286
                                                            A 19991105
                                           US 2000-201382
                                                            P 20000502
     The title compds. R4PZNR1R2 [I; R1, R2 = alkyl; R1R2 together form
AB
     alkylene; Z = alkylene; P = a direct bond, X, Y, W, XY, YW, XYW (wherein
W
     = O, S, NR3; R3 = H, alkyl; Y = UV; V = a direct bond, alkylene; U = CS,
     CO, SO2, C(:NR); R = H, OH, alkyl; X = O, NR6; R6 = H, alkyl, alkenyl,
     etc.); R4 = alkyl, alkenyl, alkynyl, etc.], useful in the activation of
     sol. guanylate cyclase, were prepd. E.g., synthesis of the urea II,
     starting with 4-bromoaniline and 1-(3-aminopropyl)pyrrolidine, was given.
     Biol. data for compds. I (e.g., IC50 for inhibition of platelet
     aggregation) were presented.
     32897-26-0P 338980-63-5P
ΙT
     RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic
     preparation); THU (Therapeutic use); BIOL (Biological study);
     PREP (Preparation); USES (Uses)
        (prepn. of amides and ureas as activators of sol. guanylate cyclase)
RN
     32897-26-0 CAPLUS
```

CN Urea, N-[3-(dimethylamino)propyl]-N'-ethyl- (9CI) (CA INDEX NAME)

EtNH-C-NH-(CH₂)₃-NMe₂

RN 338980-63-5 CAPLUS

CN Urea, N-[3-(dimethylamino)propyl]-N'-(1-pyrenylmethyl)- (9CI) (CA INDEX NAME)

REFERENCE COUNT:

REFERENCE(S):

1986,

(8) Catanese, B; BOLLETTINO CHIMICO FARMACEUTICO

V125(7), P228 CAPLUS

(9) Farbenfabriken Bayer Ag; DE 890958 C CAPLUS

(10) Glen, R; WO 0027394 A 2000 CAPLUS

(12) Hoechst Marion Roussel de Gmbh; EP 0908456 A

1999

CAPLUS

(13) Hoechst Marion Roussel de Gmbh; DE 19756388 A

1999 CAPLUS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L33 ANSWER 2 OF 14 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

2000:725451 CAPLUS

DOCUMENT NUMBER:

133:286497

TITLE:

Immunomodulatory compositions and methods of use

thereof

INVENTOR(S):

Onderdonk, Andrew B.; Tzianabos, Arthur O.; Miller,

Robert J.; Calias, Pericles

PATENT ASSIGNEE(S):

Genzyme Corporations, USA

SOURCE:

PCT Int. Appl., 62 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO. KI					ND	DATE			A	PPLI	CATI	ON N	0.	DATE					
WO 2000059490			A.	2	2000	1012		W	20	00-U	S908	7	20000406						
	WO 2000059490			90	A.	3	2001	0215											
		W:	AE,	AG,	AL,	AM,	ΑT,	AU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CR,	
			CU,	CZ,	DE,	DK,	DM,	DZ,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	GM,	HR,	HU,	

ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU,

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LV, MA, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE,
              SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZW,
              AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
          RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF,
              CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
PRIORITY APPLN. INFO.:
                                           US 1999-128177 P 19990406
OTHER SOURCE(S):
                           MARPAT 133:286497
      The invention relates to immunomodulatory compns. and related methods.
      The immunomodulatory compns. are useful for the prevention of sepsis and
      the treatment and prevention of diseases assocd. with inflammation and/or
      NOS. CM-cellulose/N-ethyl-N'-(3-dimethylaminopropyl)urea formulations
are
     described.
      32897-26-0 121007-41-8
      RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
         (immunomodulatory compns.)
RN
     32897-26-0 CAPLUS
CN
     Urea, N-[3-(dimethylamino)propyl]-N'-ethyl- (9CI) (CA INDEX NAME)
      0
EtNH-C-NH-(CH<sub>2</sub>)<sub>3</sub>-NMe<sub>2</sub>
RN
     121007-41-8 CAPLUS
     1-Propanaminium, 3-[[(ethylamino)carbonyl]amino]-N,N,N-trimethyl-, iodide
     (9CI) (CA INDEX NAME)
EtNH-C-NH- (CH_2)_3 - N+Me_3
           • I-
L33 ANSWER 3 OF 14 CAPLUS COPYRIGHT 2001 ACS
ACCESSION NUMBER:
                          2000:368337 CAPLUS
DOCUMENT NUMBER:
                           133:4656
TITLE:
                           Preparation of heteroarylpyrazoles as p38 kinase
                          inhibitors
INVENTOR(S):
                          Anantanarayan, Ashok; Clare, Michael; Collins, Paul
                          W.; Crich, Joyce Z.; Devraj, Rajesh; Flynn, Daniel
L.;
                          Geng, Lifeng; Graneto, Matthew J.; Hanau, Cathleen
E.;
                          Hanson, Gunnar J.; Hartmann, Susan J.; Hepperle,
                          Michael; Huang, He; Khanna, Ish K.; Koszyk, Francis
                          J.; Liao, Shuyuan; Metz, Suzanne; Partis, Richard A.;
                          Perry, Thao D.; Rao, Shashidhar N.; Selness, Shaun
```

Raj; South, Michael S.; Stealey, Michael A.; et al.

PATENT ASSIGNEE(S):

G.D. Searle & Co., USA

SOURCE:

PCT Int. Appl., 1210 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent English

LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

	PATENT NO.					ND	DATE			Ą	PPLI	CATI	ON N	ο.	DATE			
	WO 2000031063				A1 20000602				WO 1999-US26007						19991117			
		W:					AU,											
							EE,											
			IN,	IS,	JP,	KE,	KG,	ΚP,	KR,	KΖ,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	MD,
							MX,											
							TT,			UG,	US,	UZ,	VN,	YU,	ZA,	ZW,	AM,	AZ,
							RU,											
		RW:	GH,	GM,	KE,	LS,	MW,	SD,	SL,	SZ,	TZ,	UG,	ZW,	ΑT,	BE,	CH,	CY,	DE,
							GB,								SE,	BF,	ВJ,	CF,
2270	D = m-					GA,	GN,	GW,										
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		URCE									, , .					_		
AB	Tit	те с	ompa	5 • [.	L; R.	1 =	H, O	H, N	H2,	(cyc.	10) a.	тк (е:	u) ÀT	, ac	λт,	aryı,	, et	c.; R2
		l, hai	10,	атку.	L, a.	rkox	Α, (un) si	upst:	itute	ea p	ıper:	ıdın	ут,	etc.	; R3	=	
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$$Me_{2}N-CH_{2}-CH_{2}-NH-C-NH-(CH_{2})_{3}$$

CN Urea, N-[3-(dimethylamino)propyl]-N'-[3-[5-(4-fluorophenyl)-4-(4-pyridinyl)-1H-pyrazol-3-yl]propyl]- (9CI) (CA INDEX NAME)

REFERENCE COUNT:

10

REFERENCE(S):

(1) Anantanarayan, A; WO 9852937 A 1998 CAPLUS

(2) Anantanarayan, A; WO 9852940 A 1998 CAPLUS

(3) Fujisawa Pharmaceutical Co; EP 0531901 A 1993

CAPLUS

(4) Lilly Co Eli; EP 0846687 A 1998 CAPLUS

(5) Oku Teruo; WO 9419350 A 1994 CAPLUS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L33 ANSWER 4 OF 14 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

2000:307141 CAPLUS

DOCUMENT NUMBER:

132:331676

TITLE:

Fluorescence immunoassays using analyte

(analog)-conjugated porphyrin-silicon complex
fluorescent dyes free of aggregation and serum

binding

INVENTOR(S):

Devlin, Robert Francis; Dandliker, Walter Beach;

Arrhenius, Peter Olaf Gustaf

PATENT ASSIGNEE(S):

Hyperion, Inc., USA

SOURCE:

U.S., 58 pp., Cont.-in-part of U.S. 5,880,287.

CODEN: USXXAM

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 6060598 US 5403928 US 5641878 US 5677199 US 5880287	A A A A A A INFO.:	DATE 20000509 19950404 19970624 19971014 19990309	US 1997-874820 US 1991-701449 US 1994-333603 US 1994-346098 US 1995-476544 US 1990-523601 B2 US 1990-524212 B2 US 1991-701449 A3	19970613 19910515 19941102 19941129 19950606 19900515 19900515
			US 1994-333603 A2 US 1994-346098 A2	19910515 19941102 19941129 19950606

AB Fluorescence immunoassay methods are provided which use fluorescent dyes which are free of aggregation and serum binding. Such immunoassay methods

are thus, particularly useful for the assay of biol. fluids, such as serum, plasma, whole blood and urine. The compds. of the invention, whose

prepn. is described, include silicon complexes with porphyrin derivs. which are linked to an analyte or analog thereof, e.g. a caged dicarboxy silicon phthalocyanine digoxin probe.

IT 267422-47-9P

RL: ARG (Analytical reagent use); BUU (Biological use, unclassified); PRP (Properties); SPN (Synthetic preparation); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); PREP (Preparation); USES (Uses)

(fluorescence immunoassays using analyte (analog)-conjugated porphyrin-silicon complex fluorescent dyes free of aggregation and serum binding)

RN 267422-47-9 CAPLUS

CN Poly(oxy-1,2-ethanediyl), .alpha.-hydro-.omega.-methoxy-, ester with hydrogen (OC-6-13)-[3-[[[(5.beta.,12.beta.,14.beta.)-21,23-epoxy-12,14-dihydroxy-23-oxo-24-norchol-20(22)-en-3-yl]amino]carbonyl]-29H,31H-

phthalocyanine-2-carboxylato(3-)-.kappa.N29,.kappa.N30,.kappa.N31,.kappa.N

32]bis[[2-[[[[3-[(hydroxy-.kappa.O)dimethylsilyl]propyl]amino]carbonyl]aminoll]ethyl]carbamato]silicate(1-) (2:1) (9CI) (CA INDEX NAME)

PAGE 1-A

$$\begin{array}{c|c} \text{MeO---} & \text{CH}_2\text{-}\text{CH}_2\text{-}\text{O} & \text{NH} \\ \hline & \text{C-NH-CH}_2\text{-}\text{CH}_2\text{-}\text{NH} \\ \hline & \text{C-NH-CH}_2\text{-}\text{CH}_2\text{-}\text{NH} \\ \hline \end{array}$$

09/350,193

PAGE 1-B

PAGE 1-C

PAGE 2-A

● H+

PAGE 2-B

IT 267422-48-0P 267422-49-1P 267422-50-4P 267422-51-5P 267422-52-6P 267422-53-7P 267422-54-8P

RL: ARG (Analytical reagent use); BUU (Biological use, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); PREP (Preparation); USES (Uses) (fluorescence immunoassays using analyte (analog)-conjugated porphyrin-silicon complex fluorescent dyes free of aggregation and serum binding)

RN 267422-48-0 CAPLUS

CN Poly(oxy-1,2-ethanediyl), .alpha.-hydro-.omega.-methoxy-, ester with hydrogen

(OC-6-13)-[3-[[[(5.beta.,14.beta.)-21,23-epoxy-14-hydroxy-23-oxo-24-norchol-20(22)-en-3-yl]aminol]carbonyl]-29H,31H-phthalocyanine-2-carboxylato(3-)-.kappa.N29,.kappa.N30,.kappa.N31,.kappa.N32]bis[[2-[[[[3-

[(hydroxy-.kappa.O)dimethylsilyl]propyl]amino]carbonyl]aminol]ethyl]carbam ato]silicate(1-) (2:1) (9CI) (CA INDEX NAME)

PAGE 1-B

PAGE 1-C

PAGE 2-A

● H+

siladodecanoato(2-)]silicate(3-) (9CI) (CA INDEX NAME)

__/

PAGE 2-B

PAGE 1-B

PAGE 2-A

• H+

__/

PAGE 2-B

RN 267422-50-4 CAPLUS

CN Poly(oxy-1,2-ethanediy1), .alpha.-hydro-.omega.-methoxy-, 1,1'-diester with trihydrogen (OC-6-13)-[3-[[[4-(5-ethylhexahydro-2,4,6-trioxo-5-

pyrimidinyl)phenyl]amino]carbonyl]-29H,31H-phthalocyanine-2-carboxylato(3-

-)-.kappa.N29,.kappa.N30,.kappa.N31,.kappa.N32]bis[11-(hydroxy-.kappa.O)-11-methyl-6-oxo-2,5,7-triaza-11-siladodecanoato(2-)]silicate(3-) (9CI) (CA INDEX NAME)
- *** STRUCTURE DIAGRAM IS NOT AVAILABLE ***
- RN 267422-51-5 CAPLUS
- CN Poly(oxy-1,2-ethanediyl), .alpha.-hydro-.omega.-methoxy-, 1,1'-diester with trihydrogen
- (OC-6-13)-[3-[[[2-[[[4-(4-hydroxy-3,5-diiodophenoxy)-3,5-diiodophenyl]acetyl]amino]ethyl]amino]carbonyl]-29H,31H-phthalocyanine-2-carboxylato(3-)-.kappa.N29,.kappa.N30,.kappa.N31,.kappa.N32]bis[11-

(hydroxy-.kappa.O)-11-methyl-6-oxo-2,5,7-triaza-11-siladodecanoato(2-)]silicate(3-) (9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 1-B

PAGE 1-C

$$-NH-C-CH_2$$
OH
OH

PAGE 2-A

● H+

PAGE 2-B

RN 267422-52-6 CAPLUS

CN Poly(oxy-1,2-ethanediyl), .alpha.-hydro-.omega.-methoxy-, 1,1'-diester with trihydrogen

(OC-6-13)-[3-[[[2-[[4-(acetylamino)benzoyl]amino]ethyl]et hylamino]carbonyl]-29H,31H-phthalocyanine-2-carboxylato(3-)-.kappa.N29,.kappa.N30,.kappa.N31,.kappa.N32]bis[11-(hydroxy-.kappa.O)-11methyl-6-oxo-2,5,7-triaza-11-siladodecanoato(2-)]silicate(3-) (9CI) (CA INDEX NAME)

*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***

RN 267422-53-7 CAPLUS

CN Poly(oxy-1,2-ethanediyl), .alpha.-hydro-.omega.-methoxy-, 1,1'-diester with trihydrogen (OC-6-13)-[3-[[[4-(5-ethylhexahydro-4,6-dioxo-5-

pyrimidinyl)phenyl]amino]carbonyl]-29H,31H-phthalocyanine-2-carboxylato(3-

```
)-.kappa.N29,.kappa.N30,.kappa.N31,.kappa.N32]bis[11-(hydroxy-.kappa.O)-11-
     methyl-6-oxo-2,5,7-triaza-11-siladodecanoato(2-)]silicate(3-) (9CI) (CA
     INDEX NAME)
*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***
     267422-54-8 CAPLUS
     Poly(oxy-1,2-ethanediyl), .alpha.-hydro-.omega.-methoxy-, ester with
     dihydrogen (OC-6-13)-[3-[[(carboxydiphenylmethyl)amino]carbonyl]-29H, 31H-
phthalocyanine-2-carboxylato(4-)-.kappa.N29,.kappa.N30,.kappa.N31,.kappa.N
32]bis[[2-[[[[3-[(hydroxy-.kappa.0)dimethylsilyl]propyl]amino]carbonyl]ami
     no]ethyl]carbamato]silicate(2-) (2:1) (9CI) (CA INDEX NAME)
*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***
REFERENCE COUNT:
                         48
REFERENCE(S):
                         (1) Anon; WO 9118006 1981 CAPLUS
                         (2) Anon; EP 0260098 1987 CAPLUS
                         (5) Anon; JP 63264674 1988 CAPLUS
                         (6) Anon; EP 0336879 1989 CAPLUS
                         (7) Anon; WO 9002747 1990 CAPLUS
                         ALL CITATIONS AVAILABLE IN THE RE FORMAT
L33 ANSWER 5 OF 14 CAPLUS COPYRIGHT 2001 ACS
ACCESSION NUMBER:
                       1999:795794 CAPLUS
DOCUMENT NUMBER:
                         132:35701
TITLE:
                         Preparation of imidazolyl derivatives as as agonists
                         or antagonists of somatostatin receptors
                         Thurieau, Christophe Alain; Poitout, Lydie Francine;
INVENTOR(S):
                         Galcera, Marie-Odile; Gordon, Thomas D.; Morgan,
                         Barry; Moinet, Christophe Philippe
PATENT ASSIGNEE(S):
                         Societe de Conseils de Recherches et d'Applications
                         Scientifiques, S.A., Fr.
SOURCE:
                         PCT Int. Appl., 342 pp.
                         CODEN: PIXXD2
DOCUMENT TYPE:
                         Patent
LANGUAGE:
                         English
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
     PATENT NO.
                     KIND DATE
                                          APPLICATION NO.
                      A2
                            19991216
                                          WO 1999-US12760 19990608
        W: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ,
             DE, DK, EE, ES, FI, GB, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP,
            KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN,
            MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM,
            TR, TT, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD,
            RU, TJ, TM
        RW: GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW, AT, BE, CH, CY, DE, DK,
            ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG,
            CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
    AU 9944257
                      A1
                            19991230
                                          AU 1999-44257
                                                            19990608
    EP 1086086
                            20010328
                      Α1
                                          EP 1999-927323
                                                            19990608
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R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, PT, IE,

NO 2000006267 A 20010207 NO 2000-6267 20001211
PRIORITY APPLN. INFO.: US 1998-89087 P 19980612
US 1998-96431 A1 19980612
WO 1999-US12760 W 19990608

OTHER SOURCE(S): MARPAT 132:35701

The title compds. [I; R1 = H, (CH2)mCO(CH2)mZ1, (CH2)mZ1, etc.; Z1 = (un)substituted benzo[b]thiophene, Ph, naphthyl, etc.; R2 = H, alkyl; R1 and R2 taken together with the nitrogen atoms to which they are attached form II-IV; R3 = (CH2)mE(CH2)mZ2; E = O, S, CO, etc.; Z2 = H, alkyl, NH2, etc.; R4 = H, (CH2)mA1; A1 = C(:Y)NX1X2; C(:Y)X2; C(:NH)X2, X2; Y = O, S; X1 = H, alkyl, etc.; X2 = alkyl, etc.; R5 = alkyl, (un)substituted aryl, etc.; R6 = H, alkyl; R7 = alkyl, (CH2)mZ4; Z4 = (un)substituted Ph, naphthyl, indolyl, etc.; m = O-6] which are useful as agonists or antagonists of somatostatin receptors (no data), and for inhibiting the proliferation of Helicobacter pylori, were prepd. Thus, activating 2-furancarboxylic acid with carbonyldiimidazole followed by addn. of 2-{(1S)-1-amino-2-(indol-3-yl)ethyl}-4-phenyl-1H-imidazole afforded 94% the title compd. V. Compds. I are effective at 0.01-10.0 mg/kg/day.

IT 252305-00-3P 252311-37-8P 252311-82-3P

252314-08-2P 252314-32-2P

RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of imidazolyl derivs. as as agonists or antagonists of somatostatin receptors)

RN 252305-00-3 CAPLUS

CN Urea, N-[(1S)-5-amino-1-(4-phenyl-1H-imidazol-2-yl)pentyl]-N'-[3-(dimethylamino)propyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Ph
$$\sim$$
 (CH₂) 3 NMe₂

RN 252311-37-8 CAPLUS

CN Urea, N-[3-(dimethylamino)propyl]-N'-[(1R)-2-(1H-indol-3-yl)-1-(4-phenyl-1H-imidazol-2-yl)ethyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 252311-82-3 CAPLUS

CN Urea,

 $\begin{array}{lll} N-[3-(dimethylamino)propyl]-N'-[1-[4-(1,1-dimethylethyl)-1H-imidazol-2-yl]-2-(1H-indol-3-yl)ethyl]-& (9CI)& (CA INDEX NAME) \end{array}$

$$\begin{array}{c|c} H & O \\ |l & \\ NH-C-NH-(CH_2)_3-NMe_2 \\ |l & \\ N-& \\ Bu-t \\ N \end{array}$$

RN 252314-08-2 CAPLUS

CN Urea,

N-[2-(diethylamino)ethyl]-N'-[(1R)-2-(1H-indol-3-yl)-1-(4-phenyl-1H-imidazol-2-yl)ethyl]-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 252314-32-2 CAPLUS

CN Urea,

N-[2-(diethylamino)ethyl]-N'-[(1S)-2-(1H-indol-3-yl)-1-(4-phenyl-1H-imidazol-2-yl)ethyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L33 ANSWER 6 OF 14 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

1999:783937 CAPLUS

DOCUMENT NUMBER:

132:22973

TITLE:

Preparation of pyrrolo[2,3-d]pyrimidines as adenosine

receptor antagonists

INVENTOR(S):

Castelhano, Arlindo L.; McKibben, Bryan; Witter,

David

PATENT ASSIGNEE(S):

Cadus Pharmaceutical Corp., USA

SOURCE:

PCT Int. Appl., 169 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent English,

LANGUAGE: FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

```
PATENT NO.
                      KIND DATE
                                           APPLICATION NO. DATE
                                          WO 1999-US12135 19990601
                            19991209
    WO 9962518
                      A1
        W: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ,
             DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS,
             JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK,
             MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ,
             TM, TR, TT, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ,
            MD, RU, TJ, TM
        RW: GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW, AT, BE, CH, CY, DE, DK,
             ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG,
             CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
    AU 9942265
                                          AU 1999-42265
                            19991220
                                                            19990601
                      A1
                            20010206
                                           BR 1999-11612
                                                            19990601
    BR 9911612
                       Α
                                                            19990601
    EP 1082120
                                           EP 1999-926107
                            20010314
                      Α1
            AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO
    NO 2000006090
                            20010131
                                           NO 2000-6090
                                                            20001130
                     А
PRIORITY APPLN. INFO.:
                                        US 1998-87702
                                                         Ρ
                                                            19980602
                                        US 1999-123216
                                                         Ρ
                                                            19990308
                                        US 1999-126527
                                                         Ρ
                                                            19990326
                                        WO 1999-US12135 W
                                                            19990601
                        MARPAT 132:22973
OTHER SOURCE(S):
    Title compds. [I; R = NR1R2; R1-R4 = H, alkyl, aryl, etc.; NR1R2 =
```

heterocyclyl; R5,R6 = H, halo, alkyl, aryl, etc.; R4R5,R5R6 = atoms to complete a ring] were prepd. Thus, 2- amino-3-cyano-4,5-dimethyl-1-(1phenylethyl)pyrrole was N-benzoylated and the product cyclized to give, after deprotection and chlorination, I (R3 = Ph, R4 = H, R5 = R6 =

Me)(II;

R = Cl) which was aminated by trans-4-hydroxycyclohexylamine to give II

(R

= trans-4-hydroxycyclohexylamino). Data for biol. activity of I were given.

IT 251946-33-5P 251946-34-6P

RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of pyrrolo[2,3-d] pyrimidines as adenosine receptor antagonists)

RN 251946-33-5 CAPLUS

CN Urea, N-[2-[(5,6-dimethyl-2-phenyl-1H-pyrrolo[2,3-d]pyrimidin-4-yl)amino]ethyl]-N'-methyl- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & H & Me \\ & & N & N & Me \\ \hline \\ O & & N & Me \\ \hline \\ MeNH-C-NH-CH_2-CH_2-NH & Me \\ \end{array}$$

RN 251946-34-6 CAPLUS

CN Urea, N-[2-[(5,6-dimethyl-2-phenyl-1H-pyrrolo[2,3-d]pyrimidin-4-yl)amino]ethyl]-N'-ethyl- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & & H & \text{Me} \\ & & & & N & & H \\ & & & & N & & Me \\ \\ & & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & &$$

REFERENCE COUNT:

20

REFERENCE(S):

- (1) Chen Yuhpyng, L; WO 9413676 A 1994 CAPLUS(2) Ciba Geigy AG; EP 0682027 A 1995 CAPLUS
- (3) Hitchings, G; US 3037980 A 1962 CAPLUS
- (4) Hoechst India Ltd; IN 157280 A 1986 CAPLUS
- (5) Iwamura, H; J Med Chem 1983, V26, P838 CAPLUS
- ALL CITATIONS AVAILABLE IN THE RE FORMAT

L33 ANSWER 7 OF 14 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

1999:763780 CAPLUS

DOCUMENT NUMBER:

132:10496

TITLE:

Method for preparing thin liquid samples for

microscopic analysis

INVENTOR(S):

Berndt, Klaus W.

PATENT ASSIGNEE(S):

Becton, Dickinson and Company, USA

SOURCE:

Eur. Pat. Appl., 14 pp. CODEN: EPXXDW

DOCUMENT TYPE:

Patent

English LANGUAGE:

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

KIND DATE PATENT NO. APPLICATION NO. DATE _____ _____ EP 1999-108936 EP 961109 A2 19991201 19990505 EP 961109

20000719 A3

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO

JP 2000002839 A2 20000107 JP 1999-147168 19990526 A 19980527 US 1998-85851 PRIORITY APPLN. INFO.:

A method for producing thin samples of liqs. for microscopic anal. involves depositing a drop of the liq. sample onto the upper surface of a microscope slide near the center of the slide, positioning a flexible cover glass onto spacers on the slide, applying a downward force to the upper surface of the cover glass so that the lower surface of the cover glass touches the sample, suspending the application of force, and obtaining a thin liq. sample. A liq. blood sample prepd. this way had a central area A contg. plasma but no red blood cells. This region A was surrounded by a wide ring B contg. huge nos. of isolated red blood cells in a well-defined monolayer arrangement. Ring B was surrounded by an even

wider belt that contained red blood cells in Rouleaux formation where the length of the Rouleaux blocks increased with increasing distance from the center. This kind of blood sample prepn. does not result in morphol. changes as obsd. in the wedge slide method or during drying of blood films

in the open air.

154088-80-9, LaJolla Blue

RL: ARG (Analytical reagent use); DEV (Device component use); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); USES (Uses)

(deposited on microscope slide; method for prepg. thin liq. samples

for

microscopic anal.)

154088-80-9 CAPLUS RN

Poly(oxy-1,2-ethanediyl), .alpha.-hydro-.omega.-methoxy-, ether with CN dihydrogen (OC-6-12)-bis(2-hydroxyethyl 11-hydroxy-11-methyl-6-oxo-2,5,7-

triaza-11-siladodecanoato-011) [29H, 31H-phthalocyanine-2, 3-carboxylato(4-)-N29, N30, N31, N32 silicate(2-) (2:1) (9CI) (CA INDEX NAME)

*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***

L33 ANSWER 8 OF 14 CAPLUS COPYRIGHT 2001 ACS 1998:789144 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER:

130:38377

TITLE:

Preparation of heteroarylpyrazoles as p38 kinase

inhibitors

INVENTOR(S):

Anantanarayan, Ashok; Clare, Michael; Collins, Paul W.; Crich, Joyce Zuowu; Devraj, Rajesh; Flynn, Daniel L.; Geng, Lifeng; Hanson, Gunnar J.; Koszyk, Francis

J.; Liao, Shuyuan; Partis, Richard A.; Rao,

Shashidhar

N.; Selness, Shaun Raj; South, Michael S.; Stealey,

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Michael A.; Weier, Richard M.; Xu, Xiangdong; et al.
                         G.D. Searle and Co., USA; et al.
PATENT ASSIGNEE(S):
                         PCT Int. Appl., 828 pp.
SOURCE:
                         CODEN: PIXXD2
DOCUMENT TYPE:
                         Patent
                         English
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                   KIND DATE
                                          APPLICATION NO. DATE
     PATENT NO.
                     ____
                                          _____
     -----
                                        WO 1998-US10436 19980522
     WO 9852940 A1 19981126
        W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE,
             DK, EE, ES, FI, GB, GE, GH, GM, GW, HU, ID, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX,
            NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT,
            UA, UG, US, UZ, VN, YU, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES,
             FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI,
             CM, GA, GN, ML, MR, NE, SN, TD, TG
                                                            19980522
     AU 9875883
                     A1 19981211
                                          AU 1998-75883
                            19990524
                                         ZA 1998-4358
                                                            19980522
     ZA 9804358
                      Α
                                         EP 1998-923642
                           20000517
                                                            19980522
     EP 1000055
                     A1
           AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO
                            20000801
                                          BR 1998-9147
                                                            19980522
     BR 9809147
                     Α
                            20000121
                                          NO 1999-5695
                                                            19991119
     NO 9905695
                      Α
PRIORITY APPLN. INFO.:
                                        US 1997-47570
                                                         P
                                                            19970522
                                        WO 1998-US10436 W 19980522
                        MARPAT 130:38377
OTHER SOURCE(S):
     Title compds. [I; R1 = H, NH2, (cyclo)alk(en)yl, acyl, aryl, etc.; R2 =
Η,
     halo, alkyl, alkoxy, etc.; R3 = pyridyl, pyrimidinyl, quinolyl, etc.; R4
     H, alkyl, heterocyclyl, aryl, etc.] were prepd. Thus, R3CH2COMe (R3 =
     4-pyridinyl) was condensed with 3,4-F(MeO)C6H3CHO and the product
     cyclocondensed with TsNHNH2 to give title compd. II. Data for biol.
     activity of I were given.
ΙT
     216523-08-9P 216523-09-0P
     RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic
     preparation); THU (Therapeutic use); BIOL (Biological study);
     PREP (Preparation); USES (Uses)
        (prepn. of heteroarylpyrazoles as p38 kinase inhibitors)
     216523-08-9 CAPLUS
u
=> d his
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          36270 S L1 FULL
L3
                STRUCTURE UPLOADED
L4
L5
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L8
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              4 S ETHYL(L)DIMETHYLAMINOPROPYL(L)UREA
L9
             3 S L9 AND 1/NC
L10
             1 S L10 AND 1/0
L11
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L12
L13
            2 S L11/THU
   FILE 'USPATFULL' ENTERED AT 10:55:19 ON 01 JUN 2001
        2 S L11
     FILE 'MARPAT' ENTERED AT 10:56:49 ON 01 JUN 2001
L15
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L16
             0 S L9
    FILE 'BEILSTEIN' ENTERED AT 10:57:15 ON 01 JUN 2001
L17
            1 S L11
   FILE 'REGISTRY' ENTERED AT 10:57:49 ON 01 JUN 2001
L18
             STRUCTURE UPLOADED
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L23
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L24
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    FILE 'USPATFULL' ENTERED AT 11:03:35 ON 01 JUN 2001
L25
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    FILE 'REGISTRY' ENTERED AT 11:05:41 ON 01 JUN 2001
L26
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               STRUCTURE UPLOADED
L27
L28
          3145 S L27 FULL SUB=L26
L29
               STRUCTURE UPLOADED
L30
          1523 S L29 FULL SUB=L28
L31
          381 S L30 AND 1/0
L32
          331 S L31 NOT S/ELS
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L33
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COST IN U.S. DOLLARS
                                               SINCE FILE
                                                            TOTAL
                                                   ENTRY SESSION
FULL ESTIMATED COST
                                                   49.16
                                                            892.66
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)
                                              SINCE FILE
                                                            TOTAL
                                                   ENTRY SESSION
CA SUBSCRIBER PRICE
                                                   -7.06
                                                           -22.94
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STRUCTURE FILE UPDATES: 30 MAY 2001 HIGHEST RN 339046-06-9 DICTIONARY FILE UPDATES: 30 MAY 2001 HIGHEST RN 339046-06-9

TSCA INFORMATION NOW CURRENT THROUGH January 11, 2001

Please note that search-term pricing does apply when conducting SmartSELECT searches.

Structure search limits have been increased. See $\mbox{\scriptsize HELP SLIMIT}$ for details.

=> s 132 and 1/nr 2662225 1/NR L34 49 L32 AND 1/NR

=> d scan

L34 49 ANSWERS REGISTRY COPYRIGHT 2001 ACS
IN Urea, N-[2-(dimethylamino)ethyl]-N'-(3-phenylpropyl)- (9CI)
MF C14 H23 N3 O

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):1

L34 49 ANSWERS REGISTRY COPYRIGHT 2001 ACS
IN Urea, N-[(diethylamino)methyl]-N'-[(2-methyl-4-pyridinyl)methyl]- (9CI)
MF C13 H22 N4 O

$$\begin{array}{c|c} \mathsf{C} & \mathsf{N} \\ \mathsf{C} & \mathsf{N} \\ \mathsf{Et}_2 \mathsf{N} - \mathsf{CH}_2 - \mathsf{NH} - \mathsf{C} - \mathsf{NH} - \mathsf{CH}_2 \end{array}$$
 Me

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

=> file caplus
COST IN U.S. DOLLARS
SINCE FILE TOTAL
ENTRY SESSION
4.11 896.77

DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)
SINCE FILE TOTAL
ENTRY SESSION

CA SUBSCRIBER PRICE

0.00 -22.94

FILE 'CAPLUS' ENTERED AT 11:16:35 ON 01 JUN 2001 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2001 AMERICAN CHEMICAL SOCIETY (ACS)

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FILE COVERS 1947 - 1 Jun 2001 VOL 134 ISS 23 FILE LAST UPDATED: 30 May 2001 (20010530/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

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L13
             2 S L11/THU
     FILE 'USPATFULL' ENTERED AT 10:55:19 ON 01 JUN 2001
L14
              2 S L11
     FILE 'MARPAT' ENTERED AT 10:56:49 ON 01 JUN 2001
              0 S L11
L15
              0 S L9
L16
     FILE 'BEILSTEIN' ENTERED AT 10:57:15 ON 01 JUN 2001
L17
              1 S L11
     FILE 'REGISTRY' ENTERED AT 10:57:49 ON 01 JUN 2001
L18
                STRUCTURE UPLOADED
L19
          11247 S L18 FULL SUB=L3
L20
             50 S L18
              0 S L18 CSS
L21
              8 S L18 CSS FULL
L22
     FILE 'CAPLUS' ENTERED AT 11:01:43 ON 01 JUN 2001
L23
        10 S L22
L24
             0 S L22/THU
     FILE 'USPATFULL' ENTERED AT 11:03:35 ON 01 JUN 2001
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             7 S L22
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L26
          11247 S L18 FULL
L27
               STRUCTURE UPLOADED
L28
          3145 S L27 FULL SUB=L26
L29
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           381 S L30 AND 1/0
L31
L32
           331 S L31 NOT S/ELS
     FILE 'CAPLUS' ENTERED AT 11:14:57 ON 01 JUN 2001
L33
            14 S L32/THU
     FILE 'REGISTRY' ENTERED AT 11:16:11 ON 01 JUN 2001
L34
            49 S L32 AND 1/NR
     FILE 'CAPLUS' ENTERED AT 11:16:35 ON 01 JUN 2001
=> s 134
L35
          31 L34
=> s 134/thu
            31 L34
        375285 THU/RL
            0 L34/THU
L36
                (L34 (L) THU/RL)
=> del 136 y
=> s 135 not py>=199
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19362000 PY>=199

L36 0 L35 NOT PY>=199 => s 135 not py>=1999 2091101 PY>=1999 26 L35 NOT PY>=1999 1.37 => d ibib ab hitstr 1-26 L37 ANSWER 1 OF 26 CAPLUS COPYRIGHT 2001 ACS ACCESSION NUMBER: 1998:220203 CAPLUS DOCUMENT NUMBER: 129:4517 TITLE: Solid phase organic synthesis of polyamine derivatives and initial biological evaluation of their antitumoral activity AUTHOR(S): Tomasi, Sophie; Le Roch, Myriam; Renault, Jacques; Corbel, Jean-Charles; Uriac, Philippe; Carboni, Bertrand; Moncoq, Damien; Martin, Benedicte; Delcros, Jean-Guy CORPORATE SOURCE: Pharmacochimie de Molecules de Synthese et de Produits Naturels, Fac. de Pharmacie, Rennes, 35043, Fr. SOURCE: Bioorg. Med. Chem. Lett. (1998), 8(6), 635-640 CODEN: BMCLE8; ISSN: 0960-894X PUBLISHER: Elsevier Science Ltd. DOCUMENT TYPE: Journal LANGUAGE: English A series of N1-monosubstituted putrescine and spermine derivs. was synthesized using a solid phase methodol. Their cytotoxicity, calmodulin antagonism and polyamine uptake inhibition, pharmacol. properties shared by some antitumoral agents was evaluated. 207501-42-6P RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation) (solid phase org. synthesis of polyamine derivs. and initial biol. evaluation of antitumoral activity) RN 207501-42-6 CAPLUS CN N-[3-[4-(3-aminopropyl)amino]butyl]amino]propyl]-N'-(phenylmethyl)-, trifluoroacetate (9CI) (CA INDEX NAME) CM CRN 207501-41-5 CMF C18 H33 N5 O

O || Ph-CH₂-NH-C-NH-(CH₂)₃-NH-(CH₂)₄-NH-(CH₂)₃-NH₂ CRN 76-05-1 CMF C2 H F3 O2

F-C-CO₂H

L37 ANSWER 2 OF 26 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

1997:366218 CAPLUS

DOCUMENT NUMBER:

127:95010

TITLE:

Selective synthesis of polyamine derivatives.

Efficient derivatization of the secondary amino group

of N-monosubstituted 1,3-diamines

AUTHOR(S):

Jentgens, Christian; Hofmann, Richard; Guggisberg,

Armin; Bienz, Stefan; Hesse, Manfred

CORPORATE SOURCE:

Organisch-Chemisches Inst., Universitat Zurich,

Zurich, CH-8057, Switz.

SOURCE:

Helv. Chim. Acta (1997), 80(3), 966-978

CODEN: HCACAV; ISSN: 0018-019X Verlag Helvetica Chimica Acta

PUBLISHER: DOCUMENT TYPE:

Journal

LANGUAGE:

English

OTHER SOURCE(S):

CASREACT 127:95010

AB N-monosubstituted 1,3-diamines were selectively functionalized at the

secondary N atom via 2-phenyl-substituted hexahydropyrimidine intermediates. Reaction of the diamines with PhCHO, followed by

treatment

with an electrophile and hydrolysis, provided the desired products with excellent selectivity and in high yields. N4,N9-bis[3-phenylprop-2-

enoyl]spermine (I), which was further converted to

N1, N12-bis[3-phenylprop-

2-enoyl]spermine by a transamidation reaction, was prepd. by this way in 82% yield from spermine. Compd. I was alternatively synthesized in 83% yield, equally from spermine, by a sequence involving intermediary

protection of the terminal amino groups.

IT 191990-75-7P

RL: SPN (Synthetic preparation); PREP (Preparation)

(prepn. of polyamines by selective derivatization of secondary amino group of monosubstituted diamines)

RN 191990-75-7 CAPLUS

CN Urea, N-[3-(methylamino)propyl]-N'-(phenylmethyl)- (9CI) (CA INDEX NAME)

O || MeNH- (CH₂)₃-NH-C-NH-CH₂-Ph

L37 ANSWER 3 OF 26 CAPLUS COPYRIGHT 2001 ACS ACCESSION NUMBER: 1996:476785 CAPLUS

DOCUMENT NUMBER:

125:142463

TITLE:

INVENTOR(S):

Carbodiimide derivatives for use in biotinylations

Takenishi, Soichiro; Suzuki, Osamu; Yokomizo,

Hirohiko; Ichihara, Tatsuo; Masuda, Gen; Shiohata,

Namiko; Komiya, Kazuko

PATENT ASSIGNEE(S):

Nisshinbo Industries, Inc., Japan

SOURCE:

Eur. Pat. Appl., 55 pp.

DOCUMENT TYPE: LANGUAGE:

Patent

English

CODEN: EPXXDW

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 718300	A1	19960626	EP 1995-309433	19951222
R: DE, FR,	GB			
JP 08176159	A2	19960709	JP 1994-335492	19941222
US 5700935	Α	19971223	US 1995-577374	19951222
US 5789588	Α	19980804	US 1997-931714	19970916
PRIORITY APPLN. INFO.	:		JP 1994-335492	19941222
			US 1995-577374	19951222

OTHER SOURCE(S):

MARPAT 125:142463

Carbodiimides W1-X-N=C=N-Y-W2-Z [W1 = aliph., (un)substituted aryl, heteroaryl, tertiary amino, quaternary ammonium; -W2-Z = quaternary ammonium; X and Y = bond, alkylene; Z = biotin-contg. group] are useful

as

labeling reagents for introducing a biotin group into a nucleic acid or a protein. Thus, cyclohexyl isocyanate was treated with Me2NC6H4NH2-4 to give the urea which was converted to the carbodiimide and treated with 6-iodohexylbiotinamide to give the quaternized deriv. I.

179540-21-7P 179540-28-4P 179540-73-9P ΙT 179540-75-1P 179540-96-6P 179541-13-0P 179541-47-0P

> RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn. of carbodiimide derivs. of biotin for use in biotinylations)

179540-21-7 CAPLUS RN

CN Urea, N-[2-(dimethylamino)ethyl]-N'-(3-pyridinylmethyl)- (9CI) (CA INDEX NAME)

$$Me_2N - CH_2 - CH_2 - NH - C - NH - CH_2$$

RN 179540-28-4 CAPLUS

CN Urea, N-[(diethylamino)methyl]-N'-[(2-methyl-4-pyridinyl)methyl]- (9CI) (CA INDEX NAME)

L37 ANSWER 19 OF 26 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 1980:169045 CAPLUS

DOCUMENT NUME ER: 92:169045

TITLE: Cosmetic composition

INVENTOR(S): Grollier, Jean Francois; Fourcadier, Chantal

PATENT ASSIGNEE(S): Oreal S. A., Fr. SOURCE: Ger. Offen., 39 pp.

CODEN: GWXXBX

Patent

DOCUMENT TYPE:

the

LANGUAGE: German

FAMILY ACC. NUM. COUNT: 3

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
DE 2924230	A1	19791220	DE 1979-2924230	19790615
DE 2924230	C2	19920213	TD 1070 17000	19780615
FR 2423437	A2	19800111	FR 1978-17899	19/80013
FR 2423137	B2	19820709	JP 1979-75560	19790615
JP 55001384	A2	19800112	JP 1979-75560	19/90013
JP 63061286	В4	19881128	GB 1979-20878	19790615
GB 2024873	A	19800116	GB 19/9-200/0	19/90013
GB 2024873	B2	19820915	Ch 1070 220020	19790615
CA 1139226	A1	19830111		19800610
US 4349202	А	19820907	00 2000	19780615
PRIORITY A: 'LN. INFO.	:			19780613
				19790614
			··· · · · · · · · · · · · · · · · ·	19790614
				19790614
				19790615
			DD 20.0 C00	19790615
			02 20 10 - 0 - 0 - 0	19790615
			FR 1979-30586	19791213
			IK 13/3-30386	12/21713

AB Hair prepns. contain .gtoreq.1 polymers

[N+R1R2(CH3, mNHCONH(CH2)mN+R3R4Z]n

2X- [i., R2, R3, R4 independently = (un)substituted satd. or unsatd. aliph. or cycloaliph., (un)substituted arylaliph., NR1R2 or NR3R4 = heterocyclyl; Z = (un)substituted alkylene or alkenylene, optionally contg. .gtoreq.1 hetero atoms, or .gtoreq.1 arylene; X- = anion of an

org.
or inorg. acid; m = 2, 3], useful as carriers for dyeing or bleaching or
as permanent wave agents or lotions for treating hair before or after
permanent waving. Thus, refluxing 0-2 mol. [Me2N(CH2)3NH]2CO with 0.2

mol
Cl(CH2,5Cl in H2O 3 h gave [[N+Me2(CH2)3NHCONH(CH2)3N+Me2(CH2)6] 2Cl-]n
(I) [70698-97-4]. A bleaching agent comprised oleic acid 20,
HOCH2CH2NH2%, oleyl alc. 12, 40% tris(hydroxyethyl)ammonium lauryl
sulfate

3, Mergital OC 30 3, Me(CH2)10CON(CH2CH2OH)2 12, I 3, Bu glycol 5, EtOH 8.5, propylene glycol 6, Triton B 0.2 g. 22% Be NH4OH 18 mL, and H2O to 100 g. This formulation (60 g) was mixed with 120 g 6% H2O2 to give a gellec liq. which can be brushed on hair, where it remained 30-45 min,

was rinsed off. The wet hair was easily smoothed and had a silky feel. Similarly for the dry hair, which was also shiny and springy.

IT 70698-92-9P

RL: PREP (Preparation)

(prepn. of, for hair prepns.)

RN 70698-92-9 CAPLUS

CN Urea, N,N'-bis[3-(dimethylamino)propyl]-, polymer with 1,4-bis(bromomethyl)benzene (9CI) (CA INDEX NAME)

CM 1

CRN 52338-87-1 CMF C11 H26 N4 O

CM 2

CRN 623-24-5 CMF 08 H8 Br2

L37 ANSWER 20 OF 26 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 19

1979:557421 CAPLUS

DOCUMENT NUMBER:

91:157421

TITLE:

Phenoxyisopropanolamines

PATENT ASSIGNEE(S):

Imperial Chemical Industries Ltd., Engl.

SOURCE: Belg., 17 pp. Addn. to Belg. 808,666.

CODEN: BEXXAL

DOCUMENT TYPE:

Patent

LANGUAGE:

French

FAMILY ACC. HUM. COUNT:

THE THE COURT COUNTY

PATENT INFO MATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
BE 87 2320	A4	19790615	BE 1978-192369	19781215
ZA 7 8(5809	A	19790926	ZA 1978-5809	19781016
AU 7 843899	A1	19800424	AU 1978-40899	19781019
AU 528332	B2	19830512		
FR 24 1 31	A 2	19790720	FR 1978-35639	19781219
FR 24.1521	B2	19831014		
PRIORITY ALPLN. INFO.	:		GB 1977-52969	19771220
AB Title compds. I	(R, R1	(same or	different) = H, halo,	OH, NH2, NO2,
cyano,				

```
alkyl, cycloalkyl, alkenyl, alkynyl, alkoxy, alkylthio, cyanoalkoxy,
     alkenylomy, alkynylomy, alkanoyl, aryl, arylomy, aralkomy; R2 = H, OH,
     CH2OH, aralkoxy; Z = C2-12 alkylene; Z1 = alkylene (max. of 6 C atoms),
     C2-6 alkyleneoxy; R3 = H, indanyl, tetralinyl, oxotetralinyl, indenyl,
     1,4-dihydronaphthyl, naphthyl, R4R5R6C6H2 [R4, R5 (same or different)
same
     as R and R1; R6 = H, NH2, dialkylamine] were prepd. by several methods
and
     are useful as .beta.-adrenergic blocking agents (no data). Thus, heating
     1-(2-c::nophenoxy)-2,3-epoxypropane with H2NCH2CH2NHCONHCH2Ph in aq. EtOH
     16 h \stackrel{\cdot}{\epsilon} 90.degree. gave I (R = 2-cyano, R1 = R2 = H, Z = CH2CH2, Z1 =
CH2,
     R3 = F(.).
     71676-11-4P
ΙT
     RL: SPH (Synthetic preparation); PREP (Preparation)
        (pr pn. of, and ring cleavage of glycidyl ethers by)
     71676-1'-4 CAPLUS
RN
     Urea, +2-aminoethyl)-N'-(phenylmethyl)- (9CI) (CA INDEX NAME)
CN
```

Ph-CH2-NH- C-NH-CH2-CH2-NH2

BEST AVAILABLE COPY

L37 ANSWEL OF 26 CAPLUS COPYRIGHT 2001 ACS 1979:478762 CAPLUS ACCESSION 1 ALLER:

DOCUMENT NU LER: 91:78762

Hair coloring agents and their application TITLE:

Grollier, Jean Francois; Monnais, Christian; Peritz, INVENTOR(S):

Lyonel

PATENT ASSI WEE(S): Oreal S. A., Fr. SOURCE: Ger. Offen., 58 pp.

CODEN: GWXXBX

DOCUMENT T Patent German LANGUAGE: FAMILY ACC. U.M. COUNT: 2

PATENT INFO MATION:

PATEN	··· · · · · · · · · · · · · · · · · ·	KIND	DATE	API	PLICATION NO.	DATE
DE 28	13	A1	19790322	DE	1978-2838878	19780906
DE 28		C2	19841011			
FR 24		A1	197 90406	FR	1977-27096	19770907
FR 24	0.	B1	19800801			
FR 24		A2	19800111	FR	1978-17900	19780615
FR 241	J ä	B2	19810529			
BE 87		A1	19790306	BE	1978-190290	19780906
GB 2U	; }	A	19790321	GB	1978-35775	19780906
GB 20	₹ ±	B2	19820303			
ES 47		A1	19790401	ES	1978-473119	19780906
JP 50	· J	A2	19790418	JP	1978-108687	19780906
JP 62	-)	B4	19870219			
BR 78		А	19790502	BR	1978-5846	19780906
AU 7°		A1	19800313	AU	1978-39599	19780906

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09/350,193
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                                                              19780906
     AT 780
                       Α
                             19820715
     AT 369
                       В
                             19830225
     CH 634
                       Α
                             19830131
                                            CH 1978-9348
                                                              19780906
     CA 111
                       Α1
                             19820309
                                            CA 1978-310830
                                                              19780907
     US 435
                                            US 1980-147330
                                                              19800506
                       Α
                             19821102
PRIORITY AL: .. INFO.:
                                         FR 1977-27096
                                                              19770907
                                         FR 1978-17900
                                                              19780615
                                                              19780906
                                         US 1978-940040
              32N+R2R4Z1-X-[I, X = halogen; R1, R2 = C1-3 alkyl; R3, R4 =
ΑB
     X- -N:
              or hydroxyalkyl; Z, Z1 = C2-20 alkylene or alkylene contg.
     C1-3 a
               H2, CH2C6H4CH2, (CH2)nZ3(CH2)n where n = 2, 3, Z3 = 0, NHCONH
     CH2CH
              In hair coloring bases, which, when applied with an oxidizing
     are u:
     soln.,
              here to the hair but cause little damage. Thus, 30 g a hair
              fluid contq. 3 wt.% I[X = Br, R1-R4 = Me, Z = (CH2)3, Z1 =
     colori
            [28728-55-4] was mixed with 30 g 6% aq. H2O2 soln. to give a
     (CH2) (
            Lich was easily applied and attached to the hair to give shiny,
     cream,
              roon-color hair.
     fluff
IT
     70698-
               ₽
     RL: El
               Preparation)
                of, for hair coloring compns.)
        (p:
     70698-
RN
                CAPLUS
     Urea,
               -bis[3-(dimethylamino)propyl]-, polymer with
CN
              romomethyl)benzene (9CI) (CA INDEX NAME)
     1,4-b:
     CM
          7
     CRN :
               -17-1
               114 0
     CMF .
                                             BEST AVAILABLE COPY
Me2N= (CH2)
              TH - C - NH - (CH<sub>2</sub>)<sub>3</sub> - NMe<sub>2</sub>
     CM
     CRN ( . 1-5
     CMF ( . 1.2
BrCH<sub>2</sub>
L37 ANSWF!
               DI 26 CAPLUS COPYRIGHT 2001 ACS
ACCESSION :
               .
                          1979:421417 CAPLUS
DOCUMENT NO
              <:
                          91:21417
TITLE:
                          Quaternary ammonium polymer salts
INVENTOR (S
                         Haase, Jaroslav; Horn, Ulrich; Berendt, Hans Ulrich
```

```
PATENT ASSI
                       Ciba-Geigy A.-G., Swed.
                       Braz. Pedido PI, 59 pp.
SOURCE:
                       CODEN: BPXXDX
                       Patent
DOCUMENT TY.
                       Portuguese
LANGUAGE:
            . COUNT:
FAMILY ACC.
PATENT INFO
              <: i :
                                        APPLICATION NO. DATE
    PATENT
                   KIND DATE
                                        ______
                                                        _____
               .... ----
                                        BR 1978-3709
                                                        19780609
                          19790220
    BR 780:
                    Α
                    A3
                          19830930
                                        CH 1977-7178
                                                        19770610
    CH 6381
    CH 638
                    В
                          19840330
    US 4241
                                       US 1978-911725
                                                        19780601
                    A
                          19810127
    DD 135
                    С
                         19790829
                                       DD 1978-205841
                                                        19780607
                    A3 19811215
    SU 890
                                       SU 1978-2629400 19780607
                                                        19780608
    NL 780
                          19781212
                                       NL 1978-6242
                    Α
                                                        19780608
                                       GB 1978-26563
                          19790104
    GB 200
                    Α
                     B2
                         19820217
    GB 20(
                                        BE 1978-188476
                                                        19780609
                          19781211
    BE 868
                     Α1
                                        SE 1978-6722
                                                        19780609
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                          19781211
                                        DK 1978-2583
                                                        19780609
    DK 78€
                     Α
                          19781211
                                       FR 1978-17373
                                                        19780609
                     A1
                          19790302
    FR 235
    FR 231
                     В1
                          19821210
                     A1
                                        ES 1978-471150
                                                         19780609
                          19790901
    ES 471
                                        AU 1978-36977
                     A1
                                                         19780609
    AU 75"
                          19791213
                     A1
                                        CA 1978-305172
                                                         19780609
                          19801125
    CA 1CI
                                        PL 1978-207517
                                                         19780609
                          19801129
                     В1
    PL 111
                     A2
                          19790116
                                        JP 1978-70290
                                                         19780610
    JP 540
              NEO.:
                                      CH 1977-7178
                                                         19770610
PRIORITY # 1
              mens with amide, ester, urea, or urethane group in the
    Ione:
              y reaction of bis(tertiary amines) having these groups with
    are :
              The polymers are useful as textile dyeing auxiliaries and as
    diha]:
             1 1 flocculating agents. Thus, 0.2 mol 1,3-bis[3-
    antin
             mino)propyl]urea and 0.2 mol 4,4'-bis(chloromethyl)biphenyl
     (dime
were
             . in 200 mL refluxing MeOH to give a quant. yield of copolymer
    heat∈∷
               66-2] with inherent viscosity 2.20 dL/g (0.5% in MeOH at
     (I)
             . . A 20% aq. soln. of I was stable.
    25. dc
     69415 --
              4 69419-46-1P
ΙT
     RL: E
                maration)
               17)
       ()
RN
    6941:
               CAPLUS
              is[3-(dimethylamino)propyl]-, polymer with
CN
    Urea,
              romethyl)bennene (9CI) (CA INDEX NAME)
    CM
     CRN
                 - 7
              ... 114 0
     CMF
```

Me₂N- (CH₂ 3 - NH- (CH₂) $3 - NMe_2$

```
Me_2N - (CH_2) - NH- (CH<sub>2</sub>)<sub>3</sub>-NMe<sub>2</sub>
```

CM

CRN (2

RN 69419 APLUS

CN Urea, [3-(dimethylamino)propyl]-, polymer with

1,2-b methyl)benzene (9CI) (CA INDEX NAME)

CM

CRN ! -1 CMF (::4 O

 $Me_2N - (CH_2) = -NH - (CH_2)_3 - NMe_2$

CM 2

CRN CMF .

CH2

L37 ANSWF: 36 CAPLUS COPYRIGHT 2001 ACS

ACCESSION: 1979:122972 CAPLUS

DOCUMENT 1 90:122972

TITLE: Polymeric quaternary ammonium salts

INVENTOR(S: Haase, Jaroslav; Horn, Ulrich; Berendt, Hans Ulrich

PATENT AS: Ciba-Geigy A.-G., Switz.

SOURCE: Ger. Offen., 62 pp.

CODEN: GWXXBX

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DOCUMENT TY
                        Patent
                        German
LANGUAGE:
                :TNUC:
FAMILY ACC.
PATENT INF( :
                ::
                                          APPLICATION NO.
                                                           DATE
                           DATE
                     KIND
    PATENT
                                          _____
                                                          _____
                ____
                                          DE 1978-2824743 19780606
                           19781221
                      A1
    DE 282
                                                           19770610
                           19830930
                                          CH 1977-7178
                      A3
    CH 635
                           19840330
                     В
    CH 63€
                                          US 1978-911725
                                                           19780601
                      Α
                           19810127
    US 427
                                          DD 1978-205841
                                                           19780607
    DD 13%
                      C
                           19790829
                                          SU 1978-2629400 19780607
                      A3
                           19811215
    SU 890
                                         NL 1978-6242
                                                           19780608
                           19781212
    NL 780
                      Α
                                          GB 1978-26563
                                                           19780608
                      Α
                           19790104
    GB 20
                      В2
                           19820217
    GB 20'.
                                          BE 1978-188476
                                                           19780609
                      Α1
                           19781211
    BE 86.
                                                           19780609
                                          SE 1978-6722
                      Α
                           11781211
    SE 78
                                                           19780609
                                          DK 1978-2583
    DK 78 '
                      Α
                           19781211
                                                           19780609
                      A1
                                          FR 1978-17373
    FR 231
                           19790302
                     В1
                           19821210
     FR 23:
                                                           19780609
                      A1
                                          ES 1978-471150
                           19790901
     ES 47.
                                          AU 1978-36977
                                                           19780609
                           19791213
                      A1
     AU 75
                           19801125
                                          CA 1978-305172
                                                           19780609
     CA 1(
                      A1
                                          PL 1978-207517
                                                           19780609
                      B1 10301129
     PL 11
                                                           19780610
                                          JP 1978-70290
                      A2
                           1.790116
     JP 54
                                                           19770610
                                       CH 1977-7178
PRIORITY I
                MFO.:
                .monium polymers are prepd. from org. dihalide, esp. arom.
     Quater
                od amino derivs. of ureas, optionally mixed with other dyes
     dihal
                al as leveling agent and retarders for the dyeing of
     and a
textiles.
                1 4,4'-bis(chloromethyl)biphenyl and 0.2 mol
     Thus,
                methylaminopropyl)urea were refluxed in 200 mL MeOH, giving a
     1,3-
                 of ammonium polymer I [69420-66-2], having inherent
     100
viscosity
                 degree., 0.5% wt./vol. in MeOH). A polyacrylonitrile
     2.20 .
fabric
                versed in 200\, mL of dyeing liq. contg. 0.01 g I, adjusted to
     (5 g)
                OH, heated 100 min at 28.degree., mixed with a soln. contg.
     pH 4
                 mixt. of 3 cationic azo dyes, dyed 60 min at 98.degree.,
     0.010
                .degree., a: washed, giving a level dyeing with excellent
     cool
     wetf:
                 63419-46-1P
     6941
IT
                 paration)
     RL:
                  . for dyebath additives and coagulating agents)
       (:
                 RPLUS
RN
     6941:
                 [3-(dimethylamino)propyl]-, polymer with
CN
     Urea,
                 comethyl)bendene (9CI) (CA INDEX NAME)
     1,4-1
     CM
                 ' - 1
     CRN
                 .1 0
     CMF
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09/350,193
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1.5
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 $Me_2N-(CH_2)$:-NH-(CH₂)3-NMe₂

CM

CRN CMF

RN

69419 APLUS

13[3-(dimethylamino)propyl]-, polymer with CNUrea,

methyl)bensene (9CI) (CA INDEX NAME) 1,2-1

CM

CRN -1 14 0 CMF (

-NH-(CH₂)₃-NMe₂Me2N-(CH2

CM

CRN CMF

CH2

26 CAPLUS COPYRIGHT 2001 ACS L37 ANSW!

1976:164630 CAPLUS ACCESSION

84:164630 DOCUMENT :

Pyridyl-substituted aminoalkyl-thioureas and ureas TITLE: Durant, Graham J.; Emmett, John C.; Ganellin, Charon

INVENTOR (S

Smith Kline and French Laboratories Ltd., Engl. PATENT AS.

U.S., 6 pp. SOURCE:

COPEH: USXXAM

DOCUMENT 1 Patent
LANGUAGE: English
FAMILY ACC. OUNT: 14
PATENT IN:

	PATE:	KIND	DATE	APPLICATION NC	
	US 39	A	19760113	US 1974-450931	19740313
	GB 13	A	.9731121	GB 1971-6352	
	ZA 72	A	7721025	ZA 1972-774	19720207
	FR 21	A A5	<u>3721020</u>	FR 1972-7170	19720302
	FR 21	B1	197 51226		
	CH 57	A	19760615	CH 1972-3381	19720308
	CH 👯	,7\	19760813	CH 1975-5534	19720308
	PL 9	A P	~7 7 0331	PL 1972-153934	
	CS 1.	P	14760629	CS 1972-1579	19720309
	US 4:	\mathcal{A}	770419	us 1975-626682	2 19751029
	FI 7.	A	761130	FI 1976-3443	19761130
	FI (В	010930		
	US 4	A	19781212 19781222 19781222	US 1978-869418	3 19780116
	DK 7	A	781222	DK 1978-5802	19781222
	DK 7	A	781222		
	DK 1	C	320712		
	FI 7.	75	.790404	FI 1979-1116	19790404
	FI 6	3	-203 31		
	FI (C	20712		
	JP 5		21022	JP 1981-81781	19810527
	JP (134	50913		
	JP [©]	£.2	320618	JP 1981-168523	3 19811020
	JP ^E	15.4	331220		
	FI E.	is .	. ₹11223	FI 1981-4156	19811223
	FI (E	50628		
		C			
PRTOI		.:"0.:		GB 1971-6352	19710309
			(GB 1971-34334	19710722
				IE 1972- 13 6	19720203
			ī	us 19 72-23045 1	19720229
			ī	US 1972-290584	19720920
			1	DK 1972-909	19720228
			•	FI 1972-580	19720303
			į	US 1973-384992	19730802
			,	JP 1973-100126	19730905
			Ţ	US 1974-450931	19740313
			ì	us 1975-560909	19750321
			1	us 1976-726356	19760924
				JP 19 77-160988	19771222
AB	R(CH)	1) mNHC ':X)			n = 0, m = 3, X = S;
	R = '			n = 1, m = 2, X =	
	3-br	idvl, R1	=, $n = 1$, $n = 1$	m = 2, X = S; R =	= 2-pyridyl, R1 = Me,
n				•	
	= 1,	(=0) wer	e epd. by t	reating R(CH2)nNH	H(CH2)mNH2 with R2NCX.
	Hydi				= 3, X $=$ S) gave I
(R1	•	·	<u>-</u> .		
•	= H).	-600 : g./	C I inhibit	histamine activ	ity.
ΙT	59 0€				

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nthetic preparation); PREP (Preparation) (10 59065 CAPLUS RN hyl-N'-[2+[(2-pyridinylmethyl)amino]ethyl]- (9CI) (CA INDEX) CN Urea. NAME) H2-CH2-NH-C-NHMe ſ F 26 CAPLUS COPYRIGHT 2001 ACS L37 ANSWE 1971:**95957** CAPLUS ACCESSION : DOCUMENT !! 80: .45**957** (Aminoalkyl) imidazoles TITLE: Durant, Graham J.; Emmett, John C.; Ganellin, Charon INVENTOR (S R.; Roe, Anthony M. Smith Kline and French Laboratories <:): PATENT ASSI , Brit., 37 pp. SOURCE: COPEN: BRXXAA Patent DOCUMENT T Enc ish LANGUAGE: FAMILY AC. COUNT: 1 PATENT INF . м: APPLICATION NO. DATE KIND DATE PATENT GB 1969-56512 19691119 **GB** 134 A 19**731**219 ghty-six imi azoles (I; R = alkyl, aryl, aralkyl; R1 = H, Salts AB shenylalkyl, imidazolylalkyl; R2 = H, alkyl, substituted alkvl, alkyl; raight chain which, in some compds., was substituted by alkyl $Q = \cdot$ n = 0-3), which are histamine receptor agonists and or a , were prepose by processes which selectively introduced antage $\ensuremath{\textit{s}}$ onto one or more N atoms. Thus, 22.85 g histamine and 40 g substi : /ldiimidazole were heated 1 hr at 100.degree. and 30 min at N,N'-cre. to give .4.4 g 5-oxo-5,6,7,8-tetrahydroimidazo[1,5-c] 110-30 which, reflued with MeI in DMF, gave 2-methyl-5-oxo-5,6,7,8pyrimi idazo[1,5-/ pyrimidinium iodide (II). Refluxing 22.8 g II tet: : . 5N HCl gav + 13.5 g 1-methyl-4-(2-amino-ethyl)imidazole ove:: dih ide. 51721-88-1F ΙT **517**21 thetic preparation); PREP (Preparation) RL: S! (pi -£) RN **51721-**CAPLUS minoethyl)-''-[2-(1H-imidazol-4-yl)ethyl]- (9CI) (CA INDEX

09/350,193

CN

Urea. NAME:

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09/350,199
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of

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\text{NH-C-NH-CH}_2 - \text{CH}_2 - \text{NH}_2
    51721
                CAPLUS
RN
               aminoethyl)-!!'-[2-(1H-imidazol-4-yl)ethyl]-,
CN
    Urea, i
               medioate (1:2) (9CI) (CA INDEX NAME)
     (2Z) - 2
     CM
         1
               37-0
     CRN
     CMF
                И5 О
                HH-C-NH-CH2-CH2-NH2
     CM
              . -7
     CRN ]
     CMF C
                -1
     CDES (
Double bo: stry as shown.
                7 26 CAPLUS COPYRIGHT 2001 ACS
L37 ANSW
                         1972:400155 CAPLUS
ACCESSION .
                         77:155
DOCUMENT ::
                         Natural and artificial bleomycins. Chemistry and
TITLE:
                         antitumor activities
                         Umezawa, Hamao
AUTHOR (S):
                         Inst. Microb. Chem., Tokyo, Japan
CORPORATE :
                         Pure Appl. Chem. (1971), 28(4), 665-80
SOURCE:
                         COLEN: PACHAS
                         Journal
DOCUMENT '. '
                         Enclish
LANGUAGE:
               amine to the fermentation medium during bleomycin production
     Addn.
                formation of a bleomycin contg. that amine and suppressed
     induce
               all other pleomycins. Thus, addn. of 360 .mu.g spermidine
     format
              1 medium c. .tg. Streptomyces verticillus produced only
     [12]
               (I) [1111: -32-8]. Since only I was formed after the addn.
```

```
'1-44-3], spermine must be converted to spermidine before
     sperm
               on. Of the 42 bleomycins synthesized, those contg. diamines
     incor
               effective against Ehrlich ascites carcinoma than those contg.
     were :
               In squamous cell carcinoma 60% of the bleomycin A2
     triam
               ] remained active 1 hr after administration because of the
     [1111]
high
               the antibiotic in the tumor; however, no activity was found in
     concn.
               Bleomycins were more rapidly inactivated in liver, kidney and
     sarcor
               in lung and skin by an enzyme not yet identified.
     splee
Enzymical
               bleomycin P? was devoid of antibacterial activity, except for
     inact
               um 607 and Salmonella enteritidis.
     Myco:
ΙT
     38693-
     RL: Si
               /nthetic preparation); PREP (Preparation)
                of)
      (p)
RN
     38690
                CAPLUS
                bis(4-amino) atyl)-, compd. with 2,4,6-trinitrophenol (1:2)
CN
     Urea,
                INDEX NAME)
     (9CI)
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     CRN
               -45-8
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              ∴ N4 O
                C-NH- (CH2): -NH2
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                5 L1
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                STRUCTURE UPLOADED
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3 L4 FULL St =L3
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               3 L5 AND 3/1
L6
               S ETHYL (L) DIMETHYL (L) AMINO (L) PROPYL (L) UREA
L7
               S UREA, "N-ENHYL-N'-(3-DIMETHYLAMINOPROPYL)-"
L8
               3 ETHYL (L) DIMETHYLAMINOPROPYL (L) UREA
L9
                ; L9 AND 1/EC
L10
               3 L10 AND 1/1
L11
               10:52:57 ON 01 JUN 2001
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     FILE
                : L11
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               ... ENTERED 7.7 10:56:49 ON 01 JUN 2001
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                  1.35 H.T FT -1.99
L36
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PASSWORD:

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* * * * * *	* * * Welcome to STN International * * * * * * * * *			
NEWS 1 NEWS 2 I NEWS 3 1 NEWS 4 1 NEWS 5 A NEWS 6 7 NEWS 7 1				
NEWS EXPREDICTION NEWS HOUR: NEWS INTE	May 23 CURRENT WINDOWS VERSION IS V6.0a, CURRENT M.CI OSH VERSION IS V5.0C (ENG) AND V5.0JB (JP), AND CURRENT MISCOVER FILE IS DATED 06 APRIL 2001 STN Operating Hours Plus Help Desk Availability General Internet Information			
NEWS LOGIC	Welcome Hanna and News Items			
NEWS PHONE NEWS WWW	Direct Dial and Telecommunication Network Access to STN CAS World Wide Web Site (general information)			
Enter NEWS specific to	llowed by the frem number or name to see news on that			
All use: agreeme: research. of comme: result i:	Elease note that this agreement limits use to scientific the for soluware development or design or implementation a ligatewayer or other similar uses is prohibited and may			
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STRUCTURE . DICTIONARY	UPDATES: 3/ MAY 2001 HIGHEST RN 339046-06-9 UPDATED: 3/ MAY 2001 HIGHEST RN 339046-06-9			
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[[[(2,- if copheny))=-
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                              cophenyl)amino]carbonyl]amino]ethyl]-N-(4-
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    Ure
               (1)-N'-(.-.aet ..:yphenyl)- (9CI)
     flu
MF
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                N4 O3
                                                 OMe
               . HH-CH- CH
                                 - C-- NH-
                WERS LA YOU TSH TO SCAN? (1):1
HOW MANY
                 REGISTRY PYRIGHT 2001 ACS
     50 /
L2
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[[[(3-x hor. henyl)amino]carbonyl]amino]propyl]-N'-(4-x1)-H-(4 hen yphenyl)- (9CI)

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09/350,19
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BEST AVAILABLE GOPT

HOW MANY: YOU TISH TO SCAN? (1):1

L2 50 /: REGIS.RY COPYRIGHT 2001 ACS

IN Pro: ,

3-[[[(3-F)] nenyl) amino] curbonyl] (1-phenylethyl) amino] -N- (2-phenyl) -N- (pcl)
MF C34.

HOW MANY: WERS I YOU ISH TO SCAN? (1):1

L2 50 7 REGIO W YRIGHT 2001 ACS

IN Urea (4-nit::.phe.:d)amino]hexyl]-N'-(phenylmethyl)- (9CI)

MF C20 1 03

HOW MANY : WERS P YOU FISH TO SCAN? (1):2

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IN Ure brom a myl - [(2-fluorophenyl)methyl]-N-[6-(2-

pyr no)he: j= :I)

MF C25 . N4 O

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IN Benz. 4-chlor.-N-[2-[(4-fluorophenyl)][(4-

nitr [] amino] arbonyl]amino]ethyl]-3-nitro- (9CI)

MF C22 1 F N5 C6

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IN Ure dimental lamina propyl]-N'-(2-phenoxyphenyl)- (9CI)

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MF C18 32

NU (CHg) NMe2

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HOW MANY N THERS DO YOU WISH TO SCAN? (1):2

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IN Hept:
N-[2-[(4-: nyl)[[(2-fluorophenyl)amino]carbonyl]amino]ethy
MF C22 1 3 02

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REA OR UREAS)

L7 L(L) DIMETHYL(L) AMINO(L) PROPYL(L) UREA

=> d scan

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IN Ure: orophenyl) -N-[3-[[[(1,1-dimethylethyl)amino]carbonyl]a

minc - (4-fluorophenyl) - (9CI)



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IN Ure chophenyl) -N-[1-[3,4-dihydro-3-(3-methylphenyl)-4-oxo-

2-qu [propy1]-N-[2-(dimethylamino)ethyl]- (9CI)

MF C29 : 03

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IN Ured 1 Lightro-4-oxo-2-quinazolinyl)propyl]-N-[2-

(dime chyl]-N'-(2-fluorophenyl)- (9CI)

MF C22 1 2

-CH2-NMe2

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L7 734 AN RECISTRY COPYRIGHT 2001 ACS
IN Urea, 4-tyclohexanediylbis(methylene)]bis[N'-[4-
(dimet )ph myl]-N-(1-ethylpropyl)-, dihydrochloride, cis- (9CI)
MF C36 P . 1 C1 H
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Relative .istry.

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-[. 1-dimethylethyl)amino]-2-hydroxypropoxy]-3-(1-IN Urea, .

oxo,

MF CIS

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CM 2

Double bond recoming as shown.

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IN Urea, A 1. (4-chlore henyl) -3,4-dihydro-4-oxo-2-

quinazz propyl]-N-(2-(dimethylamino)ethyl]-N'-(3-methylphenyl)-

(9CI)

MF C29 H32 : 02

15-CH: - MP `Cl

CHOT WALLAND THE COOK

734 AMS V. REGINTRY COPYRIGHT 2001 ACS L7

Urea, N = hexyl-N-; -(3,4-dihydro-4-oxo-3-phenyl-2-quintro', copyl]-N- 2-(dimethylamino)ethyl]- (9CI) IN

MF C28 :

CH; - HII

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Ure: 1 Chapten (1) -N'-[3-(dimethylamino)propyl]-N-(2-IN

hyd: ; Cl6 L - (9CI)

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10-CHU-OH $N1_0 + (C) \rightarrow 3 = NMe2$

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                       (Ra), COO, N(Ra)CON(Rb), N(Ra)SO2, N(Ra), O, ency or a divalent group; L is a free valency, valent hydrocarbon group which may be a like; X is oxygen, optionally oxidized and nitrogen, or an optionally substituted
     S, SO, SO2;
an optionall
     interrupted
     sulfur, opti
                       on control 7 is two hydrogen atoms, oxygen, or sulfur; that R2 and an atom on ring B may together sepd. and tested as somatostatin receptor
     divalent hyd
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     form a ring?
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                                      oyl]-N'-ethyl- (9CI) (CA INDEX NAME)
     Urea, N-[3-4
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EtNH-C-NH- (CHg)
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                                      an Kodak Company; DE 2855697 A1 CAPLUS
REFERENCE(S):
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                                     man Kodak Company; JP 54145135 A CAPLUS
                                       an Kodak Company; JP 54145135 A CAPLUS
                                       man Kodak Company; GB 2010818 A 1979 CAPLUS
                                       Photo Film Co Ltd; JP 61233741 A 1986 CAPLUS
                                       TIONS AVAILABLE IN THE RE FORMAT
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ACCESSION NUMBER: :
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                             . . .tl of carbohydrate-containing dendrimers. 5.
TITLE:
                             . · p.
                                       non of dendrimers using unprotected
                                       gates
                                       , Narayanaswamy; Stoddart, J. Fraser
AUTHOR(S):
                                       ..., Univ. Birmingham, Birmingham, B15 2TT, UK
CORPORATE SOURCE:
                                       on Lett. (1997), 38(38), 6767-6770
SOURCE:
                                 EAY; ISSN: 0040-4039
PUBLISHER:
DOCUMENT TYPE:
LANGUAGE:
                                       s have been prepd. using completely
AΒ
     Carbohydrate
                                     loying a convergent growth approach. The
                                eration dendrimers, using the amide bond the possibility of obtaining densely-packed and to resort to protecting group
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                                                                                on of biotin derivative and method for pic labeling of genes by biotin derivative
 TITLE:
                                                                      Isamu; Mukai, Tsunehiro
Isamu, Japan
i Tokkyo Koho, 5 pp.
  INVENTOR(S):
  PATENT ASSIGNEE (S)
 SOURCE:
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  DOCUMENT TYPE:
  LANGUAGE:
  FAMILY ACC. NUM. ( T:
  PATENT INFORMATIO:
PATENT NO. F APPLICATION NO. DATE

JP 07157497 20 JP 1993-330034 19931201

OTHER SOURCE(S):

AB A carbodimic on deriv. (I; R1 = C1-6 alkyl, cycloalkyl; R2 = C1-6 alkylen 3, alkyl; X- = Cl-, Br-, or I-) is prepd. A non-isotopic ell ne involves biotinylation of a DNA or RNA by reacting a L1 r is biotin deriv. having a carbodimide group I. The biotin derive is colored and can be distinguishe on labeled compds., DNA, or RNA. Thus, 260 mg biotin hydra: t labeled compds., DNA, or RNA. Thus, 260 mg biotin hydra: t labeled compds., DNA, or RNA. Thus, 260 mg biotin N-B C: at labeled compds. (0.76 g) and after 15 min, the formed provate the f
  give
                100% I [:.. = .. :
                                                                                                   R3 = R4 = Me, X = Br] (II). A single
  strand
                                                                                       is dissolved in .apprx.5 .mu.L 0.1 M boric with a soln. of the carbodiimide II (50
                                                           ?
                of DNA of Hi
                                                          3.
5h.
ac.
                acid burrer
                                                                                               Her (5 .mu.L) and the mixt. was allowed to
                .mu.g/.h. (a)
                                                                                                 To the reaction mixt. was added 10 .mu.L 5
                react at .
  М
                                                                                             OH was added to ppt. biotinylated DNA, which

    Aconha brille
    d

    was remoted
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                                                                                                  dissolved in 10 .mu.L H2O. According to
                                                                                           (260 nm), 4.5 .mu.g DNA was recovered. The
                measurer :
                                                             + /
                                                                                              28 pg/.mu.L and each soln. was spotted on a cessively reacted with a ase conjugate, NBT, and BCIP. The each spot
                recover ii s
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                nitro c
                strepto
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was detected. Let u.L by blue coloration. II was also used
                         non-isotopic rel probes in the southern hybridization method.

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ACCESSION NUMBER A:
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io; Sugawa, Satoshi; Yanagida, Atsushi
i Chemical Corp., Japan
 TITLE:
 INVENTOR(S):
PATENT ASSIGNATIONS) .
                                                                                                                                                                                        . Appl., 20 pp.
SOURCE:
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  nonspecific
                             reaction upp so thyl-3-(3-dimethyl-aminopropyl)urea,
1-cycle ul-1. (1 ethyl)urea metho-p-toluenesulfone,
dimethyl no no no le, latex-immobilized digoxin, anti-digoxin
antibodu ac EDU contg. 1-ethyl3-(3-dimethylaminopropyl)-
carbod se no dested.
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ACCESSION NUL
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DOCUMENT NUM
                                                                                    Propar on of carbodimide-containing biotin
TITLE:
                                                                                   derivatives as reagents for detecting point mutation
                                                                                    of and diagnosis of hereditary disease
INVENTOR(S):
                                                                                  Ya. ...o Isamu; Mukai, Tsunehiro
                                                                                Ya: Isamu, Japan
Up E Tokkyo Koho
PATENT ASSIG: S):
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                to tit ion x = 1x. R1 = C1-6 alkyl, cycloalkyl; R2 = C1-6 thylen 3, x = 1 alogen ion), suitable for chem.
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                 naturation is the y (1) mixing for hybridization each complime try the second and its corresponding gene
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               complime try 'mastern of a normal gene and its corresponding gene assuming the process of the mutations, (2) reacting the above bioting through the column with a solution of the column with a solution of the column with a solution of the sequence of the isolated DNA fragment.

Inghost for the case involves (1) mixing for hybridization and of a normal gene and its corresponding through the column with a solution of the column with a solution of the isolated DNA fragment.

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The column try through the column with a solution of the corresponding through the confirms that it is the column try through the confirms that it is the column with a solution of the complementary single strands as sponding gene assuming the presence of the complementary single strands as sponding gene assuming the presence of the complementary single strands as sponding gene assuming the presence of the complementary single strands as sponding gene assuming the presence of the complementary single strands as sponding gene assuming the presence of the column with a solution of the column with
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                  ing genes with a restriction enzyme. The critical will be discontinuous 
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                  colver. It is a g G-T or T-G mismatching. Thus, 260 mg in 0.5 M NaHCO3 followed by adding a so
                                                                                                                  in 0.5 M NaHCO3 followed by adding a soln.
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                                                                           1 : and recrystn. from H2O to give 227.4 mg
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            a hama. A pate it lacking erythrocyte aldolase activity by restriction on a lab indIII, resp., sepd. by a agarose electrical and its sessed by restriction enzyme Rsal into 3 DM
                                                                             jested by restriction enzyme Rsal into 3 DNA.
            Both di es en e
                                                        we ted in a hybridization buffer at 100.degree. to at 42.degree. overnight followed by
            for 'C m:
adjustin;
            the F _{\rm c} th II at 30.degree. for 30 min. DNA's were seed. The EtC _{\rm c} solved in H2O, and passed to a avidin
            the 'F
 agarose
            colum.r.
                                                         du e column with 1 mM aq. biotin to sep.
                                     Leading of the second
            II- von
                                                                               the 411 bp fragment was recovered and
                                     on with the 386th adenine replaced with
             Co1.1711
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Rill C
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TITLE:
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                                                                                 on of water-soluble 1-ethyl-3-(3-
                                                           dir
                                                                                 minopropyl) carbodiimide
INVENTO: :
                                                           1:01
                                                                                Takahiro; Odagiri, Masaki; Imanari, Makoto
PATENT
                                                           Ke.
                                                                                 Ryubun Shinyoto Kaihatsu Gijutsu Kankyu
                                                           Kur
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SOURCE:
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ne (II) in arom. hydrocarbon, then treatment
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ir
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reactio
                                mixt. A soln. of EtNCS in PhMe was teated
               th . . n. (£ 1
                                n PhMe under ice cooling over 2 h, stirred
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                                and purification of proteolytic enzymes on
                                lica supports with immobilized gramicidin S
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                                A. P.; Bogomaz, V. I.; Tugai, V. A.;
AUTHOR (...
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                        Chr.,kc
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                        А. У.
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                                ladin Inst. Biochem., Kiev, USSR
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                                3HD4; ISSN: 0201-8470
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                                nity chromatog. of proteolytic enzymes were
              d by a. :hi:
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              sin, g' : cric al
                                yde, p-benzoquinone, sol. and insol.
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                              ces of the use of N-ethyl-N'-(3-
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dir " minopropyl) carbodiimide as coupling reagent
                                      fo t
                                                 reparation of meningococcal group C
                                                  aride-tetanus toxoid conjugate as vaccine
for
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AUTHOR (.
                                                   E. C.; Speijers, G. J. A.; Lutz, B. I. G.;
                                     Pre-
                                                  al, D.; Kanhai, V.; Haagmans, B.; Derks, H.
                                     J. .
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                                                  Hal group C polysaccharide-tetanus toxoid
                     apn. of canic a
                      The real set of
                     The archiver of this reagent results in a no. of stable ages between the polysaccharide and tetanus ages in the polysaccharide component and the N- trace in the reagent) and less stable ones as a consequence of the reaction, the reagent detailed and the coupling reagent for the preprior of the coupling reagent for the preprior of the coupling reagent, the reaction products and for reagent and of the residual reactivity of the polysaccharide component of the polysaccharide component of the polysaccharide component of the
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       t . •
                                                  genic activities of the conjugate was
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antig
                      e component but in an increase of its induction of IgG antibodies to the ic activity of the polysaccharide component he antigenic activity measured in anosorbent assay using antibodies to both
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	l o: .eotides in aq	ueous media
AUTHO	i M. B.; Ivanov	skaya, M. G.; Shabarova, Z. A.
CORPO:	: $:$ $:$ $:$ $:$ $:$ $:$ $:$ $:$ $:$	nosov Moscow State Univ.,
	o. USSR	
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A B .	tes commune toligonucleot.	ides were prepd. in 85-100%
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		of EtC:N:C(CH2)3NMe2 (I) at a
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a mine	SS 3. AN EUC OF CHE 16	accing amine, 0.5-4 n for
amilic.	the Art is a second to the second	
	in 4-1 ines with pKa	> 8. Thus, condensation of 20
•	. 3 mr 4 . pH 3.5 for 5	min in the presence of 0.5 mol
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                                         US .986-838082
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01 .E
                                 TENHWNR15 [I; Ar = (substituted) furyl or
\mathbf{A}^{\cdot}
                                 tj ene; B = COR2, CONR2R3, SO2R2, SO2NR2R3,
                                 11, alkowyalkyl, alkenyl, Ph, etc.; R2R3N =
                                 B = SO21(2), CO2R2] are prepd. To a soln. of
                     . r.
                             added 2- aroyl chloride, giving 88.0% of
                                  ted with 1,1-dimethyl-2-
                             ... . .ylamine t 70.degree. to give furoate ester
                             11. 23.0%). III at 2.4 .mu.g/kg/min gave 40% inse to approterenol in anesthetized dogs,
                     .i.
                                  tria in vitro) of 7.6.
T'.
                                  Thatic progration); PREP (Preparation)
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                                  ... prepn. f. beta.-adrenergic blockers)
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                                   171-, mc: hydrochloride (9CI) (CA INDEX
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TITLE
                            :a:
                                 on of carbodiimides by a phase-transfer
AU THO
                                 uzša M.; Petnehazy, Imre; Toke, Laszlo;
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                            lan ela
                            rv. Technol. Tansz., Budapesti Muszaki
CORPOI
Eayet.
                            n | 1521, Emag.
                                 Foly. (1988), 94(6-7), 246-9
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                            do the prepa. of carbodimides by dehydration long dischloride under solid-liq.
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                            In IIId K2CC? as base and a lipophile quaternary
   C
                                   erether is generally applicable for the
     Į
                                   borismiles, but esp. useful for unsym.
                     1.31
                    J::
                                 ..t of the carbodiimides prepd. have been
                           ρÍ
                                 more stable, cryst. quaternary salt.
     i
                   <u>e</u>
I'
                 10 pt
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                             a. nesulfony. chloride under phase-transfer
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                            of the off-cathyl- (9CI) (CA INDEX NAME)
MEDIH-
                            and the Nos
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                                  ing mitrosoalkylurea derivatives and their
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                                  astivity
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A''THOF
                            maer A. ...; Fadina, L. B.; Anoshina, G. M.;
                            tol ina, W. W.; Sof'ina, Z. P.
C+ RPC!
                                 e., Free Bovsk, USSR
                                  S JRC
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C IGHT 2001 ACS

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DOCUMENT TYPE:
                          Journal
 LANGUAGE:
                         Russian
     N, N-Dimethylpropanediamine, N, N-dimethylbutanediamine, and
     N,N,N'-trimethylethylenediamine were carbamoylated with suitable alkyl
     isocyanates, the urea derivs. formed were quaternized with Me tosylate,
     and the quaternized derivs. were treated with N2O3 to give nitrosoalkyl
     urea derivs., R1R2NCONR(CH2) nN+Me3 TsO- (R = H, Me or NO, R1 = Me,
     CH2CH2Cl2, or cyclohexyl and R2 = H or NO, and n = 2-4). The antitumor
     activity and toxicity of these compds. were evaluated. Toxicity of the
     disubstituted nitrosoalkylureas in comparison with choline-like
     nitrosoalkylureas was maintained at max. tolerable dose, 10-30 mg/kg,
     while that of the trisubstituted derivs. it decreased to the max.
     tolerable dose of 250-300 mg/kg. ClCH2CH2N(NO)CONMe(CH2)2N+Me TsO-
     the highest antitumor activity at 250 mg/kg. Structure-activity
relations
     are dis ussed.
TΥ
     111681-..6-8P 112557-32-1P
     RL: RCT (Reactant); SPH (Synthetic preparation); PREP (Preparation)
        (prenn. and quaternization of)
     111681- 5-5 CAPLUS
RN
     Urea, N-[3-(dimethylamino)propyl]-N'-methyl- (9CI) (CA INDEX NAME)
CN
      0
MeNH - C - NH - (CHala - NMea
RN 112557- 1-1 CAPLUS
CN Urea, N-[4-(dimethylamino)butyl]-N'-methyl- (9CI) (CA INDEX NAME)
      0
MeNH-C-NH- 'TH2 4-HMe2
L23 ANSWER 1 OF 10 CAPLUS COPYRIGHT 2001 ACS
ACCESSION NUMBER: 1988:5307 CAPLUS
DOCUMENT NUM. ER:
                        108:5307
TITLE:
                        Preparation of carbodiimides using phase-transfer
                        catalysis
AUTHOR(S):
                        Jaszay, Zsuzsa M.; Petnehazy, Imre; Toke, Laszlo;
                        Szajani, Bela
CORPORATE SO CALL
                       Tech. Univ. Budapest, Badapest, H-1521, Hung.
SOURCE:
                       Synthesis (1987), (5), 520-3
                       COLEN: SYNTBF; ISSN: 0039-7881
DOCUMENT TYPE:
                        Journal
LANGUAGE:
                        Enclish
OTHER GOURCE(S): CACREACT 108:5307
   FRICENN (R = cyclohemy), Ph, Bu, Me, Me3C; R1 = aminoalkyl, PhCH2,
    cyclohogyl, Me30) were grepd. by dehydration of ureas with arenesulfonyl
    chlorid onder solid-liq. phase-transfer conditions with solid K2CO3 as
    hase al. It M2N·Mt3 Cl- as catalyst. The method was esp. useful for the
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09/350,193

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. unther of unsym. substituted carbodismides. The basic carbodismides
      thise converted into more stable, cryst. quaternary salts.
      11681-18-8
     is: NOT (Leactant)
        (del aration of, by arylsulfonyl chloride)
    CAPLUS

1.4a, : [-(dimethylamino)propyl]-N'-methyl- (9CI) (CA INDEX NAME)
 MeNH- "ME " NUNG-IMe2
 L23 ALBOUER OF 19 CAPLUS COPYRIGHT 2001 ACS ACCESSION NO 4 2: 1937:32335 CAPLUS
 DOCUM. TT HU THE
                         11:32335
 TITLES
                         Lisrosoalkylureas based on alkylammonium salts and
                         their antitumor activity
AUTHO. 3):
                        Belyaev, A. A.; Gopko, V. F.; Radina, L. B.;
                         Lucetolchina, N. M.; Sof'ina, Z. P.; Anoshina, G. M.;
                         Pillova, T. E.
                         True. Khim., Sverdlovsk, USSR
SCURC.:
                         ill. m.-Farm. Zh. (1986), 20(5), 532-6
                         C EN: KHFZAN; ISSN: 0023-1134
DOCUMENT TYPE
                         √ snal
LANGUL HE:
                         I. sian
     . even to the compds. the exprepd. by reaction of dimethyl(aminoethyl)amine
    that appropriate poyanate, followed by either quaternization or half of the formation. In vitro tests of neoplasm inhibition showed 2
     entries of derivs. to be the most potent. Given i.p. to mice bearing
     ors, the two mochloride form was more active and more toxic
than
     The contact many salt from . Structure activity relations are discussed.
     1.1996 - 1-6P 105996-17-3P
      : DFD (Synthetic progration); PREP (Preparation)
     (dimethy)
              (dimethyl. ino)ethyl]-Ni-methyl- (9CI) (CA INDEX NAME)
MeliH- .i CH2- NMe2
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                           ydroxypropylamine aryl ester derivatives
, Sheung Tsam; Matier, William L.
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                           . . rican Hospital Supply Corp., USA
SCUFC
                           Int. Appl., 70 pp.
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	Žs.	983 0831	ZA 1982-7749 AU 1982-10120	19011112
	A1	1983 0601	AU 1982-10120	19021022
	A1	1983 1116	EP 1982-903569	19821028
		986 1210		19021028
	in, se,	GB, LI, LU,	NL. SE	
	EQ.	19870618	AU 1983-10120	19821028
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	£.	~ 9030 7	US 1986-838082	19860310
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		Ī	WO 1982-US1536	19821028
ΑF	13.00	Į	US 1986-838082	19860310
Wr.		H OH) CH2NH-	X-R1 [R = (un)substi	tuted aryl,
	-	lkylene; I	R1 = NR2COR3, NR2CON	R3R4, NR2SO2R3,
a	, , , ,	. ч = п, а.	ERVI, alkoxyalkyl, c	ycloalkyl,
W.	್ರಾರ್ೈ೨ ಂದಿನ⊏ಚಿತ್ರ	r, aralkyl;	NR3R4 = 5-7 membere	d heterocycle]

0 : • • •

: DtrAc with H2NCH2CMe2NH2 to give 57.4% A \cdots \cdots A· H 1 13 1 3 d with glycidol to give 2-FC6H4CO2R5 (R5 =mas treated with I to give . .2CC6H4F-4 (II). At 2.7 mg/kg II 3 h after nibition of heart rate response to *...* of this invention were also useful in the a-91 e · 41...c · data). IT. mounte : ration); PREP (Preparation) RHUS CII 1.52 nethyl- (901) (CA INDEX NAME)

Mewall 2000 MHz

 $\mathbf{L}^{\gamma}\mathbb{R}$ F. SYRIGHT 2001 ACS A: ...)69 CAPLUS DC : ... - .09 \mathbf{T}^{n} . ituted 3- -: _ 112.21... propanols $\mathbf{I} \cap \mathcal{F}_{i}$ Suson, Bill Bemjamin Rudolf; Hedberg, Sven re; Lundgren, Bo Torsten PATEII AB, Swed. Si: TY Pat. Appl., 25 pp. BAXXDU D. L. . . 5*** · : $\mathbf{F}_{\mathbf{r}}$ P . .

> $K^{*}\Pi$ APPLICATION NO. DATE -----706 GB 1982-35707 19821215 50807 J810 EP 1982-850257 19821210 2430 , GB, IT, LI, LU, NL, SE 10515 AT 1982-85025 AT 1982-850257 19821210 30928 ZA 1982-9249 19821215 30618 FI 1982-4339 19821216 500 NO 1982-4237 19821216 11922 1102 1701 JP 1982-219367 19821216 49216 ES 1982-518268 19821216 0428 HU 1982-4066 19821216 - 1127 CA 1982-417848 19821216 . 31 RO 1982-109344 19821216 (... CS 1982-8725 19821216

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                                               US 1985-757763
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                                            SE 1981-7574
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                                            EP 1982-850257
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                                            US 1982-450006
                                                                 19821215
                                            US 1983-482266
                                                                 19830405
                                            US 1984-621147
                                                                 19840618
 Αŀ
                  ..ol
                                    h = H, alkyl, cycloalkyl, cycloalkylalkyl; R1
                   : F2
                                   active acyl group; R3, R4 = H, acyl,
                                   A = H, alkyl, hydroxyalkyl, alkoxyalkyl;
N
                                  : Hovascular agents (no data), were prepd. by
                    . .8€
                                   4-(2-methoxyethoxy)phenyl ether was treated
                   eth
                                   olinecarboxamide to give I (R-R4 = H, n = 2,
                   .ino .
ΙT
                     - - - ;
                     ·· ;
                                  idyl aryl ethers)
\mathbb{R}^{\mathbb{I}}
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                                  -: hyl- (9CI) (CA INDEX NAME)
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                                  ropoxyphenyl derivatives and pharmaceutical
                                   ions containing them
Ι...
                                   . Richard; Louis, William John
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                                   ..-G., Switz.
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                                   . Appl., 57 pp.
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                                             EP 1981-810439
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                                           CA 1981-389517
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                                           AU 1981-77171
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                                        EP 1981-810439
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                                        US 1990-584306
                                                            19900917
                                        US 1991-782791
                                                            19911021
Al-
                  . 11
                               " loalkyl, alkenyl, cycloalkylalkyl,
                  .:d
                                   aralkenyl; R1 = H, substituent; R2 = H, R;
                                T = 0, S; Z = 0, n = 2, 3; Z = bond, n = 0
                  Ω.
1- :
                                 alky., and Q1 = bond, the R1 =
su:
                                and colgrable derivs. in esterified form, in
                                 ...ful as cardioselective .beta.-
                                were prepd. E.g., 4-PhCH2OC6H4OH was
                                our pluethyl ether, and the resultant
                                be a way thorey) benzene was debenzylated by
                  - (:.-
                                4-(2 cyclopropylmethoxyethoxy)phenol was
                  5.
                                rated with CuCN to give 2-benzyloxy-5-(2-
                  .111111
                                mitri e. The latter was debenzylated by
                  Lon
                                 Th epithlorohydrin to give
                   ant
                                 propylmethoxyethoxy) benzonitrile. The
                   1000
                   : .
                                notable.)-3-phenylurea to give
                                 care thoughphenoxy] -3-[2-(3-
                                  II was an effective
                  .
                  15
                                Ca. 82 examples of I were prepd.
                  Ele .
                                  ar taga
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                  ∍.:t.
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promary benzene derivs.) RNCN thy (CA INDEX NAME) Μ∈ . A 4 1. 101 ACS AC^{i} 4 THIES DO. T1lmulbitors of nucleotide biosynthesis. 1. re in mucleosides. 2 AU . my, and A.; Thomas, H. Jeanette; Brockman, R. Wil oz, Glynn P. **C**C. . Tell per Lab., South. Res. Inst., Birmingham,),, 17 12 m. (1981), 24(2), 184-9 10 A: (1988): 0022-2623 SO! DC^{-1} \mathbf{L}^{x} Me, as yclohexyl; R1 and R2 = H or NO; R3 = 1, ascil-1-yl; R4 = H or OH) were prepd. AŁ. 01 the idity compared to the known nitrosourea preparation); PREP (Preparation)
property (preparation) CE thr (CI) (CA INDEX NAME) Me⁻ I^{a} on , PRIE (Preparation) RM Ci: with 2,4,6-trinitrophenol (1:1)